Nano-Biosensors: Probing Intracellular Response to Nanoparticle Therapy

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For my Lily --
You can have it all!
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“Why am I discouraged? Why is my heart so sad? I will put my hope in God! I will praise him again — my Savior and my God!”

Psalm 42:5
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ABSTRACT

Nanoparticle based cellular therapies hold great promise for clinical applications and medicinal use in human patients by allowing for targeted delivery of a personalized medicine payload to specific cells and tissues in a variety of disease states. The large surface to volume ratio of solid inorganic nanoparticles and the availability of facile surface functionalization chemistries with gold, gold-coated, and semi-conductor nanoparticles enables the design of delivery agents which can simultaneously carry a targeting molecule, such as an antibody or cell penetrating peptide (CPP), a short oligonucleotide for RNA interference or intracellular sensing, and a full gene for genetic therapy to correct aberrant protein function or cause apoptosis of cancer cells. The scope of applicability for nano-therapy technology is incalculable, however current understanding of the intracellular uptake and processing of nanomaterials-based therapeutics is limited and many facets of the cellular response to nanoparticle therapy are still in need of investigation. The goal of this dissertation work has been to elucidate the effects of nanoparticle-based therapeutics to ascertain the intracellular fate and processing of model nanomedicines by designing nanoparticle-bioconjugates capable of spatiotemporally reporting live intracellular uptake and processing events using fluorescence microscopy, and magnetic detection. The first chapter gives an introduction to nanoparticles and their use in biological applications, as well as detailing how they can be used as intracellular sensors. The second chapter investigates the ability to control therapeutic DNA cargo release from a gold nanoparticle in live cells using different appendage chemistries. The third chapter probes the intracellular environment experienced by the nanotherapeutic and discusses nano-induced effects to the intracellular environment. The fourth chapter investigates nanotherapy cellular uptake targeting using cell penetrating peptides, as probed by fluorescent quantum dots, to determine outcomes for a variety of naïve and drug resistant mammalian cell lines, include human lung, skin and brain cancers as well as rodent cancer model cell lines. The fifth chapter combines optical and magnetic analysis techniques to investigate multiplexed sensing with gold shelled iron oxide nanoparticles to investigate cellular uptake. And the last chapter summarizes the work and provides a discussion of the outlook for the work.
CHAPTER 1
INTRODUCTION

1.1 Background of Nanomaterials for Biological Applications

Since the beginning of time, humans have been motivated, both by curiosity and the need for survival, to learn from and manipulate the world around them. Through this, mankind has honed the skills needed to recognize patterns, design tools, and concoct medicines to improve quality of life and expand the collective human intellect. Today those same skills are used in the pursuit of scientific insights. Driven by the fundamental desire to more deeply understand the physical world and to put that knowledge to practical use, scientific inquiry has led to great advances in knowledge, technology, and medicine.8,9 One of the most noteworthy fields of advancement, which spans all three categories, is the development and study of nanomaterials for biological and medical application.

Figure 1.1 Comparative scale of common biologically relevant objects, molecules, and nanoparticles, versus punctuation.

Materials fabricated at the nanometer scale have unique size-dependent physical properties and have the added advantage of being similar in size to biologically relevant objects and molecules (Figure 1.1). The term “nano” is used as the shorthand for the unit nanometer,
where nano- is the Greek prefix that denotes a unit one that is billion times smaller than the root unit of meter, when referring to size. Objects on the nanometer scale are at least 100 times smaller than the width of a human hair (~100 µm), and in the field of nanoscience, it is convention that a true nanoparticle is between 1-100 nm in diameter. This makes nanomaterials similar in size to biological macromolecules such as proteins and nucleic acids, as well as small cell structures, like ribosomes, while still being smaller than the smallest animal organelle, the mitochondria (~500 nm) (Figure 1.1). Added to the utility of the nanometer size regime, is the unique collection of physical phenomenon that occur in materials solely at the nanometer size scale. These properties include unique interaction with electromagnetic radiation, in the form of enhanced photoluminescence, absorption, and magnetization -- depending on the material; as well as otherwise uncommon shape morphologies (not seen in the bulk-scale material) that can allow for catalytic activity and reactive surfaces for appending molecules of biological interest. Nanosized materials also have large surface to volume ratios which provide optimum area for functionalization with biologically useful molecules. For these reasons nanometer sized materials are particularly valuable for interacting with individual living cells and directly delivering therapeutic or sensing agents for treatments and diagnostic purposes (Figure 1.2).

Figure 1.2 (A) General schematic of multi-biofunctionalized nanoparticle (NP) for diagnostic and therapeutic cellular applications with appended peptide for cellular targeting, and diagnostic or therapeutic fluorescent dye-labelled drug and nucleic acid molecules. (B) General pathway of NP cellular uptake and processing, showing endosomal encapsulation, processing, escape and disassembly of the NP construct to recover dye fluorescence, and potential exocytosis.
Gaining better understanding and more control over nanomaterials’ surface functionalization chemistry, as well as insights into cellular uptake and processing will provide new mechanisms for specifically tailoring nanomaterials for individualized therapeutic approaches. Eventually leading to the design of materials with the ability to diagnose intracellular conditions as indicators of disease, deliver therapeutic cargo to treat or reprogram unhealthy cells, or target and specifically destroy nuisance cells, like cancers. The overarching goal of bio-nanomaterials research is to develop and deploy an army of “smart” nano-medicines which are able to infiltrate cells of interest, reconnoiter health status, and complete therapeutic missions to benefit patients, at the cellular level. A major benefit of the eventual realization of these nanotherapeutic technologies will be a cessation of the dreaded pharmaceutical “side-effects” that are so prevalent in modern medicinal approaches which use small molecule drugs that are systemically delivered to circulate the whole body, as they often have numerous non-target interactions, causing cascading needs for additional drugs to combat unwanted symptoms.\textsuperscript{16,17} Thus, the ideal utilization of nanomedicine will take healthcare to the next level of sophistication and improve quality of life for all mankind. The first step to actualizing this utopic goal is to fully understand the physical and biological implications of these nanomaterials, as well as to recognize what tools are available to control and optimize their efficacy.

1.1.1 Nanomaterials: History, Composition, and Physical Properties

Millions of years before humans could fathom designing or using nanomaterials, nature had engineered nanometer-sized structures into the scales of butterfly wings, the pads of lizard feet, and the surfaces of lotus leaves and shark skin, for their optical, adhesive, and antifouling/antimicrobial properties.\textsuperscript{18,19} In the last few thousand years humans took note of substances that possessed these advantageous properties and began to inadvertently use nanomaterials in decorative paints, the coloring of glass, and as homeopathic elixirs, due to their vivid colors and healing effects.\textsuperscript{20} It has only been in the last century that scientific technology has advanced to the state that nanoparticles can thoroughly be investigated.

The Chemist and 1925 Nobel Prize in Chemistry winner, Richard Adolf Zsigmondy, was the first person to be able to understand the mechanism of how gold salts, when heated, incorporated into molten glass to yield red “ruby” or “cranberry” glass, which gains its signature
color from the gold nanoparticles suspended in the glass matrix. Zsigmondy was able to
determine that small colloids of gold were responsible for the red color of “ruby” glass though
analysis using the ultramicroscopy technique he developed, where incident light is scattered by
the nanoparticles embedded in the glass. This was one of the first demonstrations of the ability to
classify materials at the nanoscale. And evidence that nanometer sized materials could
possess vastly different properties from the bulk and atomic material, as bulk gold and ionic gold
salt are both golden-yellow in color, in contrast to the gold nanoparticles which clearly interacted
with light in a much different way to yield its brilliant red appearance.

Another stunning and ancient example of colloidal nanoparticles providing unique
aesthetics to glass can be seen in the Lycurgus Cup (Figure 1.3A) which was made with gold
and silver NP-embedded glass around the fourth century; the combination of these metallic
nanoparticles give the appearance of green glass when illuminated from the outside but
iridescent red glass when illuminated from within. The reason metallic nanoparticles (those
composed of gold, silver, copper, etc.) can have such striking iridescence and vivid coloring is
because at the nanoscale these metals develop a phenomenal property called local surface
plasmon resonance (LSPR), where the large number of surface metal atoms that loosely hold
their valence electrons, relative to the number of internal core metal atoms of the nanoparticle,
gives rise to a so called “sea of electrons” that freely oscillate in an electromagnetic field and
generate these unique interactions with visible light (Figure 1.3B). Complimentarily, the LSPR
of the metal nanoparticles also create unique light absorption profiles for each metal particle,
giving a signature plasmonic peak with some ability to discern even the approximate size of the
particle based on breadth of the peaks relative to its maxima when plotted as a function of the
molar absorptivity (ε), also known as the extinction coefficient (Figure 1.3C). This is due to
the resonant frequency of the plasmon being determined by the elemental composition of metal
nanoparticle, but the intensity being related to the overall number of aligned surface atoms able
to resonate constructively in the electromagnetic field, which decreases with decreasing particle
size due to the curvature of the surface becoming more severe as well as there being a larger
proportion of surface defects present at smaller nanoparticle sizes.

A more complete understanding of such insights has only recently been made possible
due to advances in technology and the instrumentation necessary to observe these physical
Figure 1.3 (A) Nano-gold and -silver embedded glass of the Lycurgus Cup (c. 4th century AD) under room light (i) and illuminated from within (ii) showing the unique optical properties of metal nanoparticles (Image: Public access from Trustees of the British Museum). (B) Schematic of the local surface plasmon resonance of a metallic nanoparticle depicting the sea of electrons oscillating in the electromagnetic field. And (C) visible light absorbance spectroscopy of gold nanoparticles with radii of 8.5 nm (light grey), 4 nm (dark grey), 3.3 nm (pink), 2.15 nm (red), 1.5 nm (green), and 0.95 nm (gold), showing the optical effect of NP size, plotted as molar extinction coefficient versus wavelength.
phenomenon, only seen in nanomaterials, and to test and validate the theory behind their unique behavior. The most important analytical technique to allow nanomaterials research to be validated has been electron microscopy, and particularly transmission electron microscopy (TEM), invented in 1931 by Max Knoll and Ernst Ruska.\textsuperscript{27} TEM allows for imaging of nanomaterials down to 2 nm in diameter, for elements with a large Z-number (or the number of protons in the nucleus) which determines how much X-ray radiation will be reflected to produce the negative image of nanoscale materials for the determination of their size, shape and elemental composition. Similarly, dynamic light scattering (DLS) allows for detection of the hydrodynamic radii of the NP and the surface charge can be measured by collecting the zeta-potential, allowing for more information to be collected about the nanoparticle including its hydrocarbon surface coating ligands (responsible for much of the functionality as well as the colloidal stability of the nanomaterial), which are not visible to TEM due to their low Z-number. Despite the discovery of the leading principle of DLS, Brownian motion, in 1920, the commercial instrumentation was not available until after the invention of the laser in the 1960s, which allowed for precise scattering of light calculations to be performed.\textsuperscript{28–30} In addition to the innovation of the analytical tools needed to probe the physical properties of nanomaterials, interest in the investigation of nanomaterials and nano-specific phenomenon for the development of new technologies and medicines was ignited by a series of scientific philosophers in the mid-to-late twentieth century. The first and most noteworthy of which was Richard Feynman, when in 1959, he gave his historic invitation entitled “There’s Plenty of Room at the Bottom” which challenged scientists with the notation that useful nanometer-sized structures and machines could be purposefully created through the direct manipulation of atoms. He also went a step further and charged innovators to translate this philosophical discourse into reality by promising a large monetary prize from his own resources to the first person to make a rotating electric motor smaller 1/64 inch cubed or write a page of a book on a pin head, with an optimism that it could be accomplished in short order.\textsuperscript{31} Both undertakings were completed in his lifetime.\textsuperscript{32,33} Others, such as Norio Taniguchi and Eric Drexler, also caught the vision that Feynman had and helped to solidify the term “Nanotechnology” (in 1974 and 1986, respectively) to be defined as the field of research and innovation devoted to creating and utilizing materials at the nano-scale.\textsuperscript{32,34,35}

Also in the 1980s, a similarly crucial leap in the development of nanomaterials science was made with the discovery and demonstration of the quantum confinement effect experienced
by semiconductor nanocrystals, termed quantum dots, by L.E. Brus. Brus discovered the band gap energies of aqueous CdS crystallites was dependent upon the particle size, where the bulk semiconductor (Figure 1.4A) displayed a distinctively smaller band gap energy (and thus lower energy photophysical properties) as opposed to nanometer sized semiconductor crystals, which demonstrate higher energy band gaps as the particles become smaller in diameter (Figure 1.4B-D), due to the quantized energy levels being pinned by the particle size as it nears the exciton Bohr radius.

Figure 1.4 Schematic diagram of semiconductor materials and their energy levels in (A) macroscopic “bulk” semiconductor crystals with continuous energy bands. And quantized energy levels of nanometer sized semiconductor nanocrystals showing (B) 6-4 nm diameter with red emission, (C) 4-2 nm diameter with green emission, and (D) 2-1 nm diameter with blue emission, for CdS quantum dots.
This means that the small physical parameters of the crystal lattice at the nanometer size scale cause limitations for the separation distance of an excited electron and its counterpart “hole,” thus predetermining the possible energy released by the recombination of the exciton (electron-hole pair) for a particular size of QD, making the wavelength of emitted light dependent on size. These quantum confinement effects can be explained as a three-dimensional “particle in a box” model.\textsuperscript{36} Due to the discrete energy pinning quantum confinement effect, quantum dots possess very narrow emission profiles with large absorption cross-sections.\textsuperscript{37,38} Paired with their high photostability compared to organic dye fluorophores, QDs are ideal colorful multiplexable photoluminescent agents for biological investigation, such as labels for fluorescent microscopy and as indicators in diagnostic applications.\textsuperscript{39-41}

In addition to metallic and semiconducting nanocrystals, which possess unique optical interactions with electromagnetic radiation, investigation of nanoscale magnetic materials, such as iron oxide, nickel, and cobalt, have led to great innovation in the utility of nanomaterials for diverse applications in medicine and technology.\textsuperscript{42} The physical founding of magnetic materials results from the atoms of magnetic materials possessing unpaired valence electrons which generates a net “spin” direction, or polarity, for the electrons of each atom, known as the magnetic dipole moment.\textsuperscript{43} For bulk magnetic materials, the larger number of atoms in the solid crystal lattice allows for the formation of multiple regions or “domains” of atoms that align their magnetic dipole moments in the same direction, in the absence of an external magnetic field. The ability to form multiple magnetic domains is due to the majority of magnetic materials possessing a magnetic “single-domain” size in the nanometer size-regime, which is much smaller than the bulk crystal.\textsuperscript{44} However, all bulk magnetic materials possess some degree of disorder, thus there is non-uniform orientation of the multiple magnetic domains, with some fraction of the magnetic dipole moments partially or completely cancelling one another out, and the net effect is diminished total magnetic moment observed for the material (Figure 1.5). Thus, magnetic effects can be enhanced for nanoparticles of the single-domain size for the magnetic material, over the bulk, due to the unity of magnetic spins aligning in a single direction. One such example of magnetic enhancement at the nanoscale is the generation of superparamagnetic iron oxide (SPIO) nanoparticles, where due to the quantum confinement of their surface spins, they are able to respond to a magnetic field, despite net magnetic moment not being observed in the bulk material (\textit{i.e.} rust).\textsuperscript{45}
For the abovementioned reasons, the prevalence of nanoparticle use has recently exploded, not only for the study of their unique physical properties, but also for the creation of enhanced materials, as well as for diagnostic and therapeutic biomedical applications.

1.1.2 Nanomaterials for Biological Sensing & Therapeutic Applications

Since the first herbal remedy for an illness was prepared in ancient times, the medicinal community has been relentlessly in search of more effective compounds, drugs, and therapy agents to treat the specific ailments of patients. In modern times this effort has been rebranded as a search for “personalized medicine,” which would treat each illness of each patient on an individual level. The goal of which being to forego problematic secondary interactions of therapeutic molecules, or their necessary emulsifiers and binding agents, with non-target biological elements which give rise to the dreaded “side effects” for which current pharmaceuticals are infamous. In most cases the majority of modern pharmaceutical side effects can be minor, such as binding a non-target receptor and causing headache, nausea, etc.; but in some severe cases, especially those where an allergic reaction occurs, the side effects can
be life threatening anaphylaxis, paralysis, or irreversible neurological damage.\textsuperscript{46} Thus, the need for innovation of new biomedical treatment and diagnostic approaches is highly needed.

Currently, many forms of biomedically interesting nanometer sized particles exist, from both natural and anthropogenic origins, including viral particles made primarily of nucleic acids and proteins, and synthesized nano-aggregates composed of lipids, polymers, biomolecules or combinations thereof.\textsuperscript{47–49} However, the field of synthetic inorganic nanoparticles (composed of metals, metal-oxides and semiconductor nanocrystals) holds numerous biological and commercial advantages over these other systems, which do not possess the same threshold of engineered control over therapeutic or diagnostic utility, and lack the toolbelt of tunable physical properties that inorganic nanomaterials possess.\textsuperscript{15} Examples of these include the enhanced optical and electronic properties of metal nanoparticles, sharp photoluminescent properties of semiconductor quantum dots made biologically useful with a zinc sulfide capping layer, and superparamagnetic iron oxide (SPIO) nanomaterials which can be made biologically compatible with an inert gold shell. These nanomaterials not only easily bind biologically relevant molecules to their surface for efficient therapeutic delivery, but also each possess unique visualization and analysis capabilities, not available in other delivery platforms, which allow them to be superlative biological sensors and probes of intracellular nano-therapeutic delivery.

Due to the strong plasmonic properties of gold nanoparticles (AuNP) they are not only able to absorb large amounts of energy which has applications in directed ration therapy\textsuperscript{50} and cavitation\textsuperscript{51} \textit{via} local heating for anti-cancer therapies, but AuNPs are also able to energetically couple to fluorescent dipoles (\textit{i.e.} fluorophores) and non-radiatively absorb the energy of their excited state relaxations, effectively “quenching” their fluorescence at distances described by Surface Energy Transfer (SET) theory (Figure 1.6)\textsuperscript{26,52,53} expressed by Eqn. 1.

\[
E_{SET} = \frac{1}{1+(d/d_o)^4} \quad \text{(Equation 1)}
\]

where \(E\) is the efficiency of Surface Energy Transfer (with 1 signifying 100\% quenching and 0 signifying unquenched dye emission), \(d\) is the distance between the AuNP surface and fluorophore, and \(d_o\) is the distance at which point 50\% of the fluorophore’s energy is transferred to the AuNP non-radiatively.
Application of this theory allows for dye labelling of therapeutic cargo attached to the surface of an AuNP in order to monitor cargo release as well as to design and assess aptamers for binding molecules of biological analytes, in complex environments.\textsuperscript{54-57} Additionally, the use of multiple fluorophores with different photoluminescent profiles allows for the creation of coordinated sensing responses to analytes and events using Multicolor SET (McSET),\textsuperscript{56,58} which can be used to dynamically track the delivery of differentially coupled short DNA and linearized plasmids into human cancer cells. The use of SET and McSET AuNP cellular nano-deliver agents provides information about the release rate dynamics and spatial location of the therapeutic cargo during the entire transfection process when monitored by fluorescence microscopy or similar techniques.

**Figure 1.6** Schematic representation of Surface Energy Transfer, where the 6 nm AuNP is able to non-radiatively quench emission from the excited fluorophore (fluorescein amidite: FAM) appended to a DNA molecule at the 35\textsuperscript{th} base pair away from the synthetic 5’ thiol attachment to the AuNP surface, a distance specifically chosen to correlate with the D\textsubscript{0} of 160 Å for the AuNP\textsubscript{6nm}–FITC pair.

When dye labelling of the therapeutic cargo is not feasible, due to potential chemical interactions or loss of therapeutic efficacy, the ability to use a luminescent nanoparticle delivery agent, such as a semiconductor QD, is invaluable. Additionally, a complimentary non-radiative energy transfer technique employable by QDs and other fluorophores is Förster’s Resonant Energy Transfer (FRET), which describes the coupling of two or more fluorescent dipoles, where a lower energy emitting fluorophore acts as a non-radiative energy acceptor from a higher
energy emitting donor fluorophore while the molecules are within close proximity. Thus, luminescent QDs have been used in a variety of delivery and sensing applications, as well as finding a place as diagnostic indicators.\textsuperscript{59–63}

Magnetic nanoparticles find their niche in their unique ability to act as facilitators of magnetic transfection and cell sorting, along with being useful for anti-cancer therapy, as well as contrast agents for magnetic resonance imaging (MRI) to trace NP location and processing in a live organism.\textsuperscript{64} In the former, cells can be transfected with a magnetic nanoparticle construct and internalization of the magnetic material can be augmented by application of a gentle magnetic field to physically direct the nanoparticles into the cells of interest, and once taken-up the magnetic particles can facilitate the translocation of the entire cell which is physically drawn toward the applied magnet by the ingested magnetic particles; alternatively specific antibodies can be attached to the MNP surface to allow for sorting without internalization. This technology has applications in cell culturing and genetic transformation technologies, where traditional transfection and cell sorting techniques are not as straightforward.\textsuperscript{65} In clinical applications, cancer cells permeated with MNPs can be subjected to an oscillating magnetic field, which causes physical movement of the particles and thus damage to cancerous cells.\textsuperscript{66} Less invasive studies utilizing MNP therapeutics, where light cannot penetrate tissues or optical signals are not detectable above the background, utilize MRI to quantify uptake and identify the biodistribution and fate of nano-therapeutics.\textsuperscript{67, 68}

Thus, the creation of nanoparticle-based therapeutic, imaging, and sensing materials for biological and biomedical application holds great promise. Investigation into the fundamental interactions of nano-bioconjugates with biological systems will lead to insight and innovation in the design of better nano-therapeutic agents, leading to better healthcare options for future generations.

1.2 Characterization of Nanomaterial-Bioconjugates

In order to assess the properties and efficacy of nanomaterials for therapeutic applications, the synthesis of nanomaterial-bioconjugate agents must be followed by strict scrutiny of quality, uniformity, and physical properties as assessed by physical characterization techniques. Determining the physical properties of materials at the nanoscale present unique challenges in that their size regime is far below the threshold of human senses, and they are even
too small to be visualized at the highest magnification of a light microscope, due to being smaller than the average wavelength of visible light (~500 nm). Residing between the molecular scale and bulk matter, nanomaterials require unique approaches to determine their size, shape, and electromagnetic properties. For these reasons special characterization techniques have been developed to assess the nature of the synthesized nanomaterials and evaluate the success of the attempted bioconjugations.

1.2.1 Size and Shape Analysis

The unique properties of materials at the nanoscale are greatly dependent on the size and shape of the nanoparticle. For instance, semiconductor materials at the nanoscale are termed quantum dots (QDs) due to the differences observed in their optical properties as compared to the bulk materials which are due to quantum confinement of their semiconductor band gap. This quantum effect that takes place at the nanoscale, creates unique absorbance and emission profiles based on particle diameter, even for materials with the same chemical composition.\(^6^9\) Additionally, gold nanoparticles (AuNP) have a molar absorptivity, also known as the extinction coefficient (\(\varepsilon\)), that scales with the volume of the particle, as well as local surface plasmon resonance (LSPR) that is highly dependent on shape, with different absorbance maxima profiles for spheres \textit{versus} rods and star shapes, giving these shapes unique heating profiles under infrared (IR) radiation illumination.\(^7^0\) Gold has even been observed to show emissive behavior when restricted to the size of molecular clusters.\(^7^1\) Magnetic nanoparticles (MNP) have specific single domain sizes, which can enhance or hinder the magnetism of particles of sizes at, above, or below the single domain size.\(^4^4\) Shape variations in MNPs can cause anisotropic, or orientation-dependent, effects on the interaction of the nanomaterial with a magnetic field.\(^7^2\) The shape of particles can not only have effects on particle heating rates, and electromagnetic properties, but can also effect therapeutic and sensor cargo loading levels and cellular uptake of nanoparticles.\(^7^3,7^4\) Additionally, the uniformity of nanomaterial preparations can greatly affect the ability to collect meaningful ensemble averages from optical, magnetic, and mass based measurements in cellular and animal model experiments. Thus, accurate characterization of nanomaterials after synthesis and functionalization is paramount for application reproducibility and proper interpretation of experimental findings.
The ability to visualize matter that is smaller than the wavelength of visible light (i.e. less than 0.5 µm in diameter), is of paramount importance to the field of Nanoscience and allows for the elucidation of previously invisible phenomenon, such as the growth pattern, structure, and composition of nanomaterials at fixed points in time. In 1931 the first Transmission Electron Microscope was designed by Max Knoll and Ernst Ruska, with commercial implementation taking place by the 1940s. Transmission Electron Microscopy (TEM), allows for this visualization at the nanoscale because, rather than the typical white light illumination of a traditional light microscope, TEMs use a columnated beam of electrons to illuminate materials. Since the de Broglie wavelength of an electron beam (found by the equation: \( \lambda = \frac{h}{p} \), where \( \lambda \) is wavelength, \( h \) is Planks constant and \( p \) is the momentum of the electron wave) is \(~0.01\) nm, being three orders of magnitude smaller than visible light (500 nm), it provides near atomic resolution of nanosized structures. Additionally, the principles of diffraction can be applied to assess the elemental identity of the sample using electron diffraction spectroscopy (EDS), so that sample size and composition can be obtained with the same technique.

Some limitations exist for TEM, such as the preference for a large atomic z-number or multiple layers of atoms in order to provide good contrast against the background grid, typically composed of a carbon mesh. The importance of this is increased at very small particle sizes, making the effective limit of detection on most TEMs around 2-5 nm in diameter. TEM also suffers from a lack of dimensionality in the images generated, which can lead to misidentification of nanomaterial structures that are not imaged from multiple orientations, i.e. plate-like nanomaterials could be misinterpreted to be spheres, or nanoparticles with non-uniform projections may appear to have a smaller or larger average size than is truly representative. To help combat this limitation, Scanning Electron microscopy was developed in 1938 by Manfred von Ardenne. Although these techniques are powerful for NP analysis, they are still limited to visualizing elements with large nuclear proton densities that are capable of scattering the electron beam in a detectable pattern.

The physics of particle diffusion in a solution was first observed by Robert Brown in 1827, shortly before his discovery of the cellular nucleus in 1833, and has come to be known as the phenomenon of Brownian Motion. Brown observed that where larger objects tumble in a solution, they do so at a slower rate than smaller particles in the same solution (with the same
viscosity). Later Albert Einstein further developed the “Quantitative Theory of Brownian Motion” in 1905 in order to mathematically represent the nature of particulate diffusion (Eqn. 2):

\[ P = \frac{e^{-x^2/4Dt}}{2\sqrt{\pi Dt}} \]  

(Equation 2)

Here \( P \) is the probability density for a particle, moving a certain distance, \( x \), in a given time, \( t \). \( D \) is the coefficient of diffusion for the solution, and is equal to one-half the average of the squared displacement in the x-direction.

Additionally, nanomaterials passivated with coordinating ligands (biologically relevant surface coating molecules, DNA, peptides, etc.) of various size and charge profiles can also be analyzed by utilizing a dense matrix, such as agarose or polyacrylamide gel made in an isotonic buffer, and applying an electrical current across the gel field to propel the loaded bioconjugated nanomaterial of interest through the matrix gel as a function of size and relative charge. Gel electrophoresis is an indispensable biochemical analysis tool and has been used extensively in the study of DNA, RNA and proteins for more than half a century. The technique was originally developed by Oliver Smithies in 1955 to study protein size. In the case of biologically coated nanomaterials, an important characteristic that can be determined by electrophoresis is the relative retention of the NP in the gel as an indicator of covalent or dative ligand attachment, as opposed to electrostatic ligand association with the surface of the NP. A mobility limited or retained bioconjugated NP (as compared to the as-synthesized NP counterpart) is indicative of a larger total particle size, including ligands, that cannot move through the gel matrix as easily as a more compact counterpart. Since the NP core size and net charge is not expected to change under ambient conditions, the increased gel retention is attributed to the increase in size and number of biological passivating ligands. Electrostatically bound ligands, on the other hand, are not able to maintain their association with the NP surface in the presence of an electric field and thus are said to “rip” off the surface of the NP, causing a loss of aqueous solubility of the NP, and no movement beyond the loading well for the NP. Alternatively, ligands and biomolecules with limited stability may be removed by the gel matrix at an unpredictable rate causing a smear of the bio-nanocomplex with differentially decomposed complexes being represented at near continuous size intervals, this is opposed to stably bound biomolecules on a nanocomplex that
may display a broad but single band, with broadness due to inhomogeneity of the number of bound ligands (Figure 1.7).

![Figure 1.7](image)

**Figure 1.7** (A) Schematic of principles for gel electrophoresis, showing applied current from negative above the wells, to repel negatively charged molecules, to positive at the bottom to attract molecules through the gel matrix, as well as representative post-run pattern of (i) as-synthesized aqueous AuNPs with tight band, (ii) stable large appended biomolecules with single broad band, (iii) stable smaller appended biomolecules with single broad band, and (iv) unstable appendage with biomolecule removal by the gel matrix displaying a continuous smear. (B) Photograph of AuNP-bioconjugates with stable (i) for as-synthesized passivating ligand, (ii) mono-thiol, (ii) dual thiol, (iii) thiol plus amine bidentate attachment of 30mer DNA, and unstable (iv) dual amine, and (v) mono-amine attachment of 30mer DNA.

### 1.2.2 Physical Properties of Nanomaterials Assessed by Spectroscopy

As introduced previously, nanoparticles exhibit unique behavior in the presence of electromagnetic radiation, including the strong absorption profile and proximal dipole quenching from the plasmonic resonance of metal NPs, as well as the band gap size-dependent photoluminescence profiles of semiconductor quantum dot nanocrystals. For these reasons, some of the most useful characterization of nanomaterials can be accomplished using Ultraviolet-Visible (UV-Vis) absorption spectroscopy and Photoluminescence (PL) emission spectroscopy. These analytical tools have been well utilized for decades in a variety of chemical systems, with the very first spectrophotometer being invented by Arnold O. Beckman and his colleagues at National Technologies Laboratories (later the Beckman Instrument Company) in 1940,¹ and the first spectrophotofluorimeter being invented in 1956 by Robert Bowman at the National
Figure 1.8 Original schematics of (A) Beckman DU spectrophotometer c. 1954,\textsuperscript{1,2} and (B) Amnco-Bowman spectrophotofluorometer (Images: Public access from the NIH Archive.)\textsuperscript{3}
Institutes of Health. Despite the relatively simple design of these instruments, (using light, mirrors and gratings which can be seen in the schematics in Figure 1.8), UV-Vis and PL spectrophotometers can provide a wealth of information about nanoparticle solubility, size or shape uniformity, proximity to passivating ligands, and of course, concentration according to Beer’s Law (Eqn 3).

\[ A_\lambda = \varepsilon_\lambda bC \]  
\( \text{(Equation 3)} \)

Where \( A \) is the absorbance at a given wavelength (\( \lambda \)), \( \varepsilon \) is the molar absorptivity, or “extinction coefficient,” at the chosen wavelength (\( \lambda \)), \( b \) is the pathlength that the light travels (i.e. the diameter of the cuvette), and \( C \) is the concentration. This can be especially important for biologically conjugated nanomaterials as they are often prepared in small aliquots due to the high price or complexity associated with obtaining certain biological molecules.

Another useful spectroscopic method of investigation for ascertaining information about ligand binding to nanoparticle surfaces is Infrared (IR) absorption spectroscopy, which probes the vibrations of chemical bonds. The absorption of infrared (IR) radiation is most apparent in the stretching and bending vibrations of molecular bonds, making IR, or more commonly Fourier transformed (FT-IR) spectroscopy useful to determine bonding and covalent attachment of ligands and appended biomolecules to the nanoparticle surface or to attached anchoring groups that have been pre-tethered to the nanoparticle surface. Complimentarily, Nuclear Magnetic Resonance (NMR) spectroscopy measures the magnetic relaxation of the atomic nucleus after being pulsed by a radio frequency, which is affected by the electronic environment of the atoms and gives information about the atomic neighbors of a given atomic nucleus, indicating what binding environments are present. Additionally, since the rate of hydrogen exchange for deuterium in primary amines is quite rapid, it can also be ascertained that a primary nitrogen is bound to a sterically hindering NP surface if the resonance peak of a primary amine is recovered upon addition of the NP to the compound or biomolecule.

1.2.3 Magnetic Characterization

Bioconjugated nanomaterials with a magnetic dipole moment can be characterized using unique analytical technology of a Superconducting Quantum Interference Device (SQUID), developed for bulk materials as recently as the 1960s and applied to nanomaterials just in the last
few decades.\textsuperscript{45} This instrument can be used to measure the magnetic moment of a material per mass when exposed to a magnetic field, as well as thermal and magnetic field strength dependence of the nanomaterial. In this way, biological ligand binding and biocompatible gold-shelling of magnetic nanomaterials can be assessed for their effects on the magnetism and downstream applications of magneto-nano-bioconjugates.

1.3 Nanomaterials in Cellular Applications

The ability of nanometer sized materials to interact with biological systems at the cellular level is unparalleled above the molecular scale. This gives nanoparticle-based cellular diagnostic and therapeutic technologies a unique niche to allow for multiplexing of sensors, therapies and cellular manipulation tools housed in a single entity, while still being small enough to enter, probe, deliver, and alter health conditions at the individual cell level. With the near limitless potential of nanoparticle-based cellular therapeutics and sensors, several design considerations become paramount to the creation of functional probes and delivery agents. The first of which is the mode and mechanism of cellular entry, as no intracellular sensor or therapeutic agent will be effective if denied access to the intracellular space, which is the prime occupation of the plasma membrane. Thus, understanding cellular uptake and processing, as well as equipping NP agents with adequate cellular targeting and transduction tools is the most necessary and first step to successful nano-biosensor and nanotherapeutic design.

An equally important design consideration involves assessing realistic \textit{versus} idealistic treatment and sensing goals. Although a nanoparticle can theoretically accommodate numerous different types of molecules simultaneously, sensor analysis is dictated by the ability to detect and resolve the signal over the background, regardless of methodology (fluorescence, mass, magnetism, \textit{etc.}), limiting the number of analytes or intracellular conditions that can be reliably detected in a single pass. Similarly, therapeutic approaches must overcome cellular defense mechanisms to be effective. Thus, loading particles with several redundant sensor or medicinal molecules becomes advantageous to optimize detectability and efficacy, so contemplation on what specific tasks the NP agent is to perform and what limitations will be faced by the therapy or analysis approaches of the sensor is crucial.

Finally, the fate and elimination of the NP agent must be considered, as the end-goal of nanotherapeutic and diagnostic technologies predominantly envisions use in patients or live cell
samples, where bioaccumulation and material toxicity become vital factors. Thus, agents must be small enough to escape cells and cleared by the renal system, and/or non-toxic and able to be broken down and used or excreted by biochemical pathways.

### 1.3.1 Cellular Uptake and Nanomaterial Trafficking

In order for any therapeutic to be effective, the biologically active agent needs to be able to access the intracellular target that it is designed to act upon. For bio-active nanomaterials this means that the NP must not only be loaded with the therapeutic molecule but must also be equipped with agents that assist in rudimentary cellular targeting and facilitate entry into the cells requiring treatment. A few studies of peptide and protein functionalized nanomaterials have reported direct cytosolic delivery, but the mechanisms of this process are not well understood. The majority of known NP cellular-uptake mechanisms involve some form of endocytosis, where the bio-nanomaterial construction is encapsulated by a portion of the cellular plasma membrane prior to entering into the intracellular space within the endo-lysosomal pathway (Figure 1.9A). The endo-lysosomal pathway provides cells with an innate defense against pathogens, such as viruses, as well as provides a mechanism for cellular digestion of foreign materials and transport of nutrients throughout and then outside of the cell via exocytosis. As understood from decades of research into cellular biology, the typical progression and processing of exogenous material by cells (Figure 1.9B) follows the path of membrane association (which may be mediated by cell surface receptors or electrostatic interactions with sticky glycoproteins), followed by encapsulation of the foreign matter (via clathrin or caveolin mediated, or independent invagination of the phospholipid membrane). Once the transfected material is packaged within the cell it is trafficked to an early endosome (pH ~7-6), which acidifies to a late endosome (pH ~6-5) through membrane fusion events with acid and enzyme containing vesicles, and finally the endosome fuses with a lysosome (pH ~5-4.5) in order to be fully degraded. Any molecular building blocks are recycled to the endoplasmic reticulum and Golgi network and unusable molecular debris is transported out of the cell.

As one can imagine, if a therapeutic agent never leaves this pathway, there would be no way for it to interact with its cellular target and accomplish its designed function. Thus, the need for analysis and understanding of how nanomaterial therapeutics are taken into cells, processed through the endo-lysosomal pathway, and escape prior to degradation, in order to fulfill their
designed beneficial purpose, is vital to the successful design and delivery of functional and effective cellular treatments.

Figure 1.9 Cellular uptake and processing of exogenous material showing (A) different mechanisms of endocytosis (Image: adapted from Sandvig, and van Deurs 2002)\(^4\) and (B) endolysosomal maturation of encapsulated agent from uptake, transport to early endosome (EE), late endosome (LE) and fusion with lysosome (LY), with corresponding pH gradients (red: pH 7.4 to blue: pH 4.5), followed by cellular elimination via exocytosis (Image: adapted from Carnevale 2018).\(^5\)

1.3.2 Cell Penetrating Peptides & Other Transfection Agents

The delivery of therapeutics for cellular treatments that improve disease states, or performance of preliminary experimental analysis of the benefits and risks of such therapeutics, requires that the materials must first be able to incorporate into the cells of interest. This may seem like a straightforward problem to solve, but millennia of cellular evolution have made cell membranes very effective at keeping large, polar, non-nutritive molecules and compounds safely outside of the cell, to avoid disruption of cellular function. In order for the cell to take in what it needs, the membrane has developed selective and semi-selective membrane pores, ion channels, and receptor-mediated cellular eating and drinking, known as phagocytosis and macro pinocytosis, respectively.\(^86,87\) Thus, in order to achieve cellular uptake of the designed nanomaterials, some amount of trickery is required to get the cells to recognize an obviously
foreign entity as something desirable for internalization. This can be accomplished by encapsulating the nanomaterials in lipid or polymeric coatings which hide the true nature of the nanoparticles from the cell and often allow for direct membrane fusing or endocytic encapsulation, as with cationic lipid transfection agents, such as the Lipofectamine® and JetPRIME® families of compounds. These commercially available transfection agents are proprietary concoctions of multiple amine-modified long-chain lipids which bind electrostatically with the anionic phosphate head groups of the cell membrane. Alternatively, cell transfection of nanoparticles has commonly been accomplished with methods that utilize various types of blunt force, in the form of electroporation, micro-injection, and nano-guns, which are disadvantageous for reproducibility, effective transfection, and overall viability of the cells.

In recent decades the discovery of cell penetrating peptides (CPPs) isolated from short transduction sequences of viral sheath and native transporter proteins, and, later, synthetic derivative sequences, have opened up a new and potentially more effective and less damaging path for cellular transfection of nanomaterials. Here, instead of coating particles in gooey polymers or lipids, which potentially interfere with cellular processing and the intracellular delivery of the nanotherapeutics, or subjecting individual cells to undue stress using clinically non-transferrable technology, therapeutic nanoparticle complexes can be coloaded with short peptides that allow for the potential to not only provide entry into cells of interest, but also to provide selectivity with peptide moieties that have affinity for specific cell types.

Designing nanoparticle-based transfection platforms that are able to selectively target human cells of interest using CPPs to deliver therapeutic cargo, and simultaneously report the location, rates of delivery and processing, as well as intracellular conditions that the nanoparticle’s cargo encounter, is a transformative goal.

1.3.3 Gene Therapy & DNA Sensors on Nanoparticles

The human genome was fully mapped on April 14, 2003 and has opened the door to exportation of the genetic origin of disease. This combined with the discovery of small interfering RNA (siRNA) as a defense mechanism in cells to destroy specific RNA sequences prior to translation into proteins, by Andrew Fire and Craig Mello in 1998, has led to great interest as well as many novel approaches to cancer, autoimmune, and genetic disorder treatments, as well as for some degenerative and infectious diseases. A major hurdle of gene
therapy is the packaging of medicinal nucleic acids for safe delivery into the target cells, in order to correct or replace aberrant protein expression (as in the case of cystic fibrosis), or terminate cells which have lost their apoptotic response factors (as in the case of cancers). The current state of the art is to use pathogenically inactivated or engineered viral particles, which have been shown to trigger immune response for repeat doses. A better platform for genetic medicines delivery is that of inorganic nanoparticles as they have been shown to protect their sensitive cargo from degradation and provide the ability to multiplex treatment factors as well as load targeting agents on a single particle. For this reason, many researchers have devoted their efforts to helping to establish non-viral gene therapy approaches. However, since these technologies have only just started to be investigated in the last few decades, little is known about the cellular response to the delivered therapeutics or about the potential efficacy of these agents. Thus, the use of genetically encoded fluorescent proteins has become a standard model system for gene therapy, as successful transfection of the delivered gene sequence, followed by cargo escape from the endo-lysosomal pathway, and proper cellular processing of the delivered genetic information is required for fluorescent protein turn-on, the method allows for easy evaluation of success by fluorescence detection. Fluorescent proteins are unique fluorophores in that they are not comprised of a single conjugated ring system, as are most organic dyes, nor do they behave like photoluminescent semiconductors; instead their unique protein sequences form secondary structures called beta-barrels, which allow for the exclusion of water molecules from accessing the internal amino acid residues at the center of the barrel’s center (which would quench fluorescence) and simultaneously position residues in conformational alignment so that chromophore producing residues (Ser65, Tyr66, Gly67) are able to be enzymatically oxidized to form a conjugated system capable of fluorescence upon excitation and relaxation (Figure 1.10A). Additionally, after the discovery of green fluorescent protein (GFP) in the jellyfish, Aequorea victoria, development of all colors (Figure 1.10B), enhancement of brightness, and ability to detect analytes has also been made possible. In addition to genetic expression as an indicator of successful delivery, DNA sequences can also be chemically labelled with fluorophores to assess intracellular location, the cellular environment experienced by the nucleic acid cargo, and various molecular interactions, as well as probing these questions utilizing the inherent properties of the NP delivery agents to be used as sensors of the cellular condition post transfection with nanomaterial bioconjugates.
1.3.4 Analysis of Nano-Bioconjugates in Cellular Applications

Visualization of intracellular events has been made possible by the advent of the optical microscope, also known as the light microscope, which was first built by Anton van Leeuwenhoek in the late 17th century, and allows for the bending of light through series of various curved lenses (the combination of which is also known as an objective) to attain magnification up to 1000 times the original sample size. The discovery and development of this technology has revolutionized, and continues to revolutionize, the study of life at the cellular scale, a realm invisible to the human eye. Centuries after the initial design was fabricated, laudable advancements in both the physical mechanics and electronic technology of microscopy have led to engineering of phase contrast and fluorescent microscopic techniques, along with computer software improvement of image quality, as well as automation in collection and analysis. Similar to a spectrophotofluorometer, a fluorescent microscope uses laser or selected wavelength of light, bounced around mirrors and through optic to excite the fluorophores of interest, the difference in sample (i.e. complex cell instead of a clean quartz cuvette) necessitates a sharper collection angle than the 90° used in a spectrophotofluorometer, and in fact, fluorescent light is collected back along the light path using a series of cut off filters and dichroic mirrors.
The ability to detect fluorescent agents within complex cell environments allows for more in depth investigation of cellular biology through the use of cell-structure specific labelling technologies and multichannel and overlay image collection. Here the cellular nucleus and plasma membrane can be stained with a fluorescently labelled affinity probes, and cellular events can be resolved with differentially labelled fluorescent dyes and emissive QD agents (Figure 1.11B). Additionally, the use of the live-cell chamber (which provides physiological temperature, humidity and CO₂ atmosphere directly on the microscope stage), has allowed for real-time spatiotemporal profiling of live intracellular events following nanoparticle transfection to be observed. This paired with the ability to quantify relative fluorescence intensities of nano-bioconjugate materials over the course of the cell experiment has led to important discoveries concerning how cells process nano-therapeutics and given insights into therapeutic outcomes for cells exposed to nano-agents.⁵⁶,¹⁰¹,¹⁰²

Figure 1.11  (A) Schematic of an inverted confocal fluorescent microscope, showing light path from laser to cell sample and back to photomultiplier tube (PMT) detectors with digitally converted signal on computer display. (B) Representative live-cell confocal fluorescent microscope image of Chinese hamster ovary cells stained for cell membrane with AF594 dye-labelled wheat germ agglutinin (red), nucleus with DAPI cell permeable dye (blue), and showing FAM dye-labelled DNA release from AuNP after 12 h of incubation (green).
In sum, these advances have established cellular microscopy as a preeminent tool for probing cellular phenomena, especially with regards to nanotherapeutic processing, when fluorescent labelled or photoluminescent QDs are employed. However, some limitations do exist and include the potential for poor detectability of dilute fluorophores and, coincidingly, potential phototoxicity for cells exposed to excitation light for too lengthy of collection time or too frequent intervals of collection.

For high density, non-emissive nano-therapeutics \((i.e. 20\text{nm or larger AuNP and others})\), cryogenic electron microscopy (Cryo-EM) can be employed to assess nanoparticle cellular internalization and localization, where cells are processed by flash-freezing with liquid nitrogen prior to exposure to the electron beam in order to protect sensitive cell structures from immediate ablation in the beam.\(^{124}\) However, as with any fixed cell analysis technique, Cyro-EM does not allow for dynamic imaging of cellular events as they unfold.

A complimentary technique for analysis of gene expression or other whole-cell labeling methodologies include the use of flow cytometry, also known as flow cytometry assisted cell sorting (FACS), which incorporates microfluidic engineering to rapidly push suspended cells through a laser beam to quickly collect binary fluorescence information. The advantages of the technique is high throughput and large statistical sampling; however, the technique is limited by the inability to visually monitor events in cells, and by the analysis artifacts created by punctate or multichannel fluorescence experiments.\(^{125}\)

Magnetic nanomaterials have the unique ability to act as cell sorting labels and as magnetic contrast agents. In the former, cells which have internalized the magnetic materials would be able to be drawn by an external magnetic field to quickly distinguish transfected \textit{versus} non-transfected cellular populations, and transfected cells can further be quantified using a SQUID which measures the magnetic moment of a material relative to its mass. Thus, SQUID measurements can be used to convert the magnetic moment of similarly sized magnetic nanoparticles within cell pellets of uniform mass or cell number, in order to evaluate the quantity of magnetic particles with the various cell samples. Complimentarily, the use of Magnetic Resonance Imaging (MRI) enables the determination of the density of magnetic particles \textit{versus} the non-magnetic background tissue to create a “magnetic contrast agent” within live tissues or resected cell samples, thus the magnetic particles can be visualized \textit{ex vivo} or \textit{in vivo} non-invasively, and can be used to identify the fate and distribution of particles within a patient.
model or to quantify the number of particles that have been internalized by different cell or tissue samples.

Additionally, cell destructive methods such as spectroscopic analysis of cell lysates or inductively coupled mass spectrometry (ICP-MS) of dissolved cell samples exist to determine the potential nanoparticle content within transfected and collected cell populations. For spectroscopic methods, the presence of transfected or expressed fluorophores can be examined; limitations exist in signal detectability due to high levels of noise from biomolecules and potential degradation of the target fluorophores from release of damaging cellular components during the lysis process. ICP-MS of cell samples dissolved in strong acid allows for quantification of nanomaterials by atomic abundance within the cell sample to assess the level of cellular uptake of nanoparticles.\textsuperscript{126} This technique is best used for larger nanomaterials composed of a minimal number of different elements, preferably with a high z-number, to aid in signal-to-noise ratios over the carbon-heavy cellular background, combined with the relatively small number of atoms contained in a single nanoparticle, due to their small size. For the above mentioned reasons, the ICP-MS technique is ideal for solid metal nanoparticles such as gold, silver, or platinum nano-constructions.\textsuperscript{127,128}

1.4 Thesis Overview

The work described in this thesis explores the fundamental science behind nanomaterials-bioconjugation and intracellular responses to nanomaterials probes and therapeutics as proof of concept. This work takes full advantage of the molecular beacon toolbelt and the nano-specific optical properties of gold, gold shelled, and semiconducting nanomaterials as assessed by various spectroscopic and microscopic techniques. This work also capitalizes on peptide and cationic liposome transfection methodologies as a means of cellular internalization for various bioconjugated nano-probes.

Chapter two details a study of gold nanoparticles (AuNPs) used as delivery platforms for DNA oligonucleotides and genes with various appendage strategies. Investigating control over the attachment chemistry and thus the release rate of DNA therapeutic cargos to set the foundation for personalized nanomedicine. This work compares common mono-thiol and mono-amine conjugation strategies for DNA attachments to AuNPs \textit{versus} dual thiol, dual amine and thiol plus amine bidentate conjugation schemes to determine the rates of intracellular processing
and therapeutic release for each coupling method. The use of live-cell fluorescence microscopy allows for the direct visualization of the timing of release through fluorescence “turn on” of dye-labels on the DNA that are quenched by the AuNP while the DNA is bound to the particle within the nanometal’s Surface Energy Transfer (SET) diameter. The intracellular rates of DNA release from the AuNP surface in A375 human melanoma cells were found to follow the binding activity series: dual thiol > thiol plus amine > mono thiol > dual amine ≈ mono amine, for the investigated DNA- AuNP coupling routes.

Chapter three focuses on the simultaneous monitoring of the timing of cargo release from a nanotherapeutic AuNP as well as elucidating the intracellular pH experienced by the cargo during cell processing. This was accomplished using a triple dye-labeled duplex DNA appended to an AuNP. Utilizing the environmentally sensitive dye, pH-responsive fluorescein, paired with another Surface Energy Transfer coupled dye, DyLight405, the pH at and after cargo release was ratiometrically determined. Additionally, a non-energy-coupled dye, DyLight700, was used for nanoparticle tracking throughout the experiment. The Multicolor Surface Energy Transfer (McSET) beacon of the cargo uptake, release, and processing was visualized using live-cell fluorescent microscopy, and it was observed that the nanotherapeutic packaging of labeled DNA provides a protective effect against lysosomal degradation in acidic organelles as noted by cargo experiencing an average intracellular pH near 6.

Chapter four investigates the role of drug resistance status in cancer cells on the targeting and therapeutic delivery of nano-cargo, by assessing cellular uptake of fluorescent quantum dots functionalized with various cell penetrating peptides (CPPs). The QD-CPP nano-bioconjugation creates a labelled delivery package to investigate the nuances and difficulties of drug transport into multidrug resistant (MDR) cancer cells for potential future clinical applications of diverse nanoparticle therapeutic delivery strategies. In this study, eight distinct cell penetrating peptides (CAAKA, VP-22, HIV-TAT, HIV-gp41, Ku-70, and hCT(9-32), integrin-β3, and K-FGF) were used to study the different cellular uptake profiles in cancer versus drug resistant melanoma (A375 & A375-R), mesothelioma (MSTO & MSTO-R), and glioma (rat 9L & 9L-R, and human U87 & LN18), cell lines. The results demonstrate that cell penetrating peptide uptake varies with the amount of drug resistance and cell type, likely due to changes in cell surface markers. This study provides insight to developing functional nanoparticle delivery systems in drug resistant cancer models.
Chapter five contains a collection of analytical tools employable for evaluating nano-therapy in cells transfected with gold shelled magnetic nanomaterials, termed “magnetic gold particle” (MGP). These particles contain a magnetic iron oxide core, able to be evaluated for magnetic moment and resonance via SQUID and MRI, respectively; with the added advantages of a gold shell to allow for good biocompatibility, easy biofunctionalization with cell penetrating peptides, as well as plasmonic functionality via SET to visualize intracellular therapeutic delivery using fluorescent microscopy. This multiplex sensor brings the best aspects of magnetic and gold nanoparticles into one useful platform to answer questions of therapeutic targeting and uptake with cell penetrating peptides in cancer cell lines.
CHAPTER 2
SURFACE COUPLING CONTROLLED INTRACELLULAR CARGO RELEASE OF THIOL AND AMINE MODIFIED DNA FROM A GOLD NANOPARTICLE

2.1 Introduction

Research into micro- and nano-biological phenomenon has skyrocketed in the last few decades due to technological advances in instrumentation and new methodologies for cellular and molecular visualization. Some of the most advantageous innovations in this field include live-cell confocal fluorescence microscopy and Surface Energy Transfer (SET) based nanoparticle beacons.\textsuperscript{55,56,129–131} The pairing of these techniques allows for spatiotemporal tracking of intracellular phenomena in real time, such as the delivery of nano-therapeutic cargo into human and relevant-model cell lines.\textsuperscript{101,132} The ability of nanomaterials to load multiple therapeutic agents has been established for a variety of nanoparticle constructions.\textsuperscript{102,133–135} But the ability to control the timing of release of nano-cargo within a live cell – without external interventions that would not be possible in a true patient care model – has not been demonstrated. Further investigation of the intracellular processing of nano-delivered therapeutic cargo will lead to insights relevant to biomedical applications, such as controlled drug delivery and coordinated gene therapy approaches the for improved treatment and prevention of disease.

For the multiplexing of therapeutic cargo delivery from a nanomaterial to be truly advantageous, the ability to control the timing and location of delivered drugs or genes inside of target cells, such as cancers or diseased tissue, must be realized. Some studies have been performed which evaluate the use of external signals such as chemical activation or using electromagnetic radiation to heat and melt the particle to induce therapeutic cargo release,\textsuperscript{109,136,137} which while effective in cell culture complicate a realistic treatment approach in a living patient, where systemic interactions are possible and direct observation of the cells being treated is not. A more practical approach to controlled therapeutic release lies in utilizing surface appendage chemistry to control the dynamics of nanoparticle release in cellular environments. It is widely known that AuNPs bind to sulfur groups effectively,\textsuperscript{138,139} and that mono-thiol groups readily disassemble in the cellular environment.\textsuperscript{102,132,140} An earlier study revealed the release of short DNA \textit{versus} plasmid from a nanoparticle surface could be manipulated by use of a thiol linkage.\textsuperscript{132} Studies using SET probes and gold nanoparticle-DNA hybridization have also
demonstrated that greater stability is achieved by multidentate binding. The mechanism and cellular metabolites that control release of assembled cargo on a gold nanoparticle, once endocytosed into a cell, is not well understood, although cellular reducing agents are thought to play a role. Additionally, studies on the differences between various members of the most common mono- and bi-dentate thiol ligands have been conducted, predominantly coupled to bulky molecules or synthetic polymer chains to aide in colloidal stability under physiological conditions, which result is particles with large hydrodynamic radii. These modifications may cause unknown interactions with cellular processing and elimination of the nano-cargo by likely effecting release of therapy molecules, thus convoluting any cellular uptake and release studies. Correlating the intracellular dynamics for cargo release with the functional coupling chemistry for nucleic acids bound to a AuNP could enhance our understanding of the potential for nanoparticle-nucleic acid therapeutic agents.

In the current work, the control of intracellular release of nucleic acids cargoes covalently coupled through mono- and bi-dentate thiol and amine functional groups at the surface of a 6 nm AuNP is investigated. The study employs a multicolor surface energy transfer (SET) bio-optical transponder (BOT) to monitor uptake and release of a Dylight 488 (DL488)/ Dylight700 (DL700) labeled short, 39 base pair (bp) DNA, as well as a dye labelled 6 kbp plasmid designed to initiate production of fluorescent protein, tdTomato, following transcription. The nucleic acid cargoes are bound to the surface of the gold by appending a functionalized C₆ spacer from phosphate backbone (DNA-PO₃O-C₆H₁₂-NH₂ or DNA-PO₃O-C₆H₁₂-SH). Functionalization of the 3’ and complimentary 5’ end of the DNA allows incorporation of a mono-thiol (3’ -SH only), a dual-thiol (3’ -SH and comp 5’ -SH) and thiol plus amine coupling (3’ -SH and comp 5’ -NH₂). It is hypothesized that the strength of covalent attachment of the therapeutic cargo to the AuNP will follow: bidentate thiol (SS) > bidentate thiol plus amine (SN) > monodentate thiol (SX), followed by the weak coordination of bidentate and monodentate amines, (NN) and (NX). Elucidating the release kinetics of these simple coupling strategies will allow for control over the timing of therapeutic release from the AuNP surface in order to create coordinated treatments, which often rely on sequential delivery of therapeutics, especially in drug resistant cancer cells, such as melanoma. Understanding and utilizing AuNP surface chemistry that is able to produce controllable, self-terminating conjugation of biologically active cargo to the nano-delivery agent
will allow for tunable release of therapeutics in the intracellular environment. Realization of this goal advances nanotherapeutics one step closer to personalized medicine.

2.2 Materials & Methods

2.2.1 Materials

Water soluble, spherical 6.0 nm AuNP were synthesized by standard literature protocols for aqueous AuNP citrate nanoparticles.\(^\text{11}\) Briefly, a sparged aqueous solution of tetrachloroaauric acid hydrate (Strem Chemicals) was reduced in the presence of citric acid and tannic acid (Sigma) at 60°C with rapid stirring at a ratio of 1:4:0.5 mL of each 1% m/v solution in 100 mL H\(_2\)O total volume. The prepared AuNPs were exchanged with bis-(p-sulfonatophenyl) phenylphosphine (bSPP) (Sigma) as the protecting group. Stoichiometric displacement of the coordinated BSPP passivation layer was accomplished by thiol place exchange reactions using 5’ thiol terminated DNA sequences, as described previously.\(^\text{147}\) The synthetic DNA sequences were purchased from Midland Oligos and are listed below:

CATG-DL700-39mer-SH
5’- CATGCGACTGTGACAAATCTTAGCTGCGATAGAGTAGTC(C\(_6\)-SH) -3’

CATG-39mer-NH\(_2\)
5’- CATGCGACTGTGACAAATCTTAGCTGCGATAGAGTAGTC(C\(_7\)-NH\(_2\)) -3’

SH-DL488-comp
5’- (SH-C\(_6\))GACTACT(DL488)CTATCGGCAGCTAAGATTGTCACAGTCG -3’

NH\(_2\)-DL488-comp
5’- (NH\(_2\)-C\(_6\))GACTACT(DL488)CTATCGGCAGCTAAGATTGTCACAGTCG -3’

X-DL488-comp
5’- GACTACT(D4L88)CTATCGGCAGCTAAGATTGTCACAGTCG -3’

2.2.2 Preparation of AuNP-SX/SS/SN-DNA

AuNP-SX-DNA was prepared by complete thiol place exchange of the BSPP passivation layer using a 5’-C\(_6\) thiol with or without a 3’ C6 thiol or amine terminated short duplex DNA, to give the SX (mono-thiol), SS (dual-thiol), and SN (thiol and amine) linkages. The DNA consisted of a synthetic 35 bp sequence containing a 5’-C\(_6\) thiol and an internal DL488 dye label incorporated at the 10th bp and a complimentary strand with 3’-C6 thiol (SS) or amine (SN)
modification, or unmodified 3’ (SX), with an internal DL700 dye label incorporated at the 30th bp. Dyes were incorporated through internal modification of the dT base. The exchange was carried out in a 20:1 mole ratio of DNA to AuNP, in the presence of reducing agent, tris(2-carboxyethyl)phosphine (Sigma), resulting in displacement of ~20 passivating sites on the AuNP surface. The loading of the DNA was confirmed by gel electrophoresis (1% agarose) indicated a single red band that was retained stronger than the AuNP-BSPP, consistent with formation of AuNP-DNA.

2.2.3 Preparation of AuNP-TSX/TSS/TSN-plasmid constructs

The AuNP-TSX/TSS/TSN-plasmid constructs were prepared by initial assembly of a linearized plasmid, appended to the SX/SS/SN linker sequences (with DL488 and DL700 internal labeling), to the AuNP surface by thiol place exchange, in the presence of reducing agent, tris(2-carboxyethyl)phosphine (Sigma). The dye labels for each sequence were located on the 30th bp from the terminal thiol modification via an internal labeling of the T base.

The linker modified plasmid was prepared using the commercially available tdTomato-C1 plasmid (Clontech) digested with PCII (New England Biolabs (NEB)) following the manufacturer’s protocol. The plasmid contained a CMV promoter to ensure over expression of the fluorescent protein within the cell following the translation of the delivered plasmid. The digested plasmid (single cut site) was analyzed on a 1% agarose gel to validate digestion. The synthetic SX/SS/SN-DNA linker strands (35/39 bp) contained a 4 bp overhang and was ligated to the linearized plasmid by standard T4 ligation methods (NEB) to create tdTomato plasmids with protected 5’-C6 thiol modification with or without (TSX) a complimentary 3’-C6 protected thiol (TSS) or amine (TSN) modification to the phosphate backbone. The sequences are available in Supporting Information. The ligated plasmids containing the chemically protected functional spacers were precipitated by addition of EtOH and stored at 4°C. Isolation of the linearized DNA was verified by UV-Vis analysis of the 260nm absorption for DNA.

The protected thiol plasmid was appended to the AuNP through formation of an AuNP-sulfur bond (AuNP-TSX-plasmid), dual AuNP-sulfur bonds (AuNP-TSS-plasmid), or AuNP-sulfur and AuNP-amine bonds (AuNP-TSN-plasmid), following TCEP reduction (20 mM, 2 h, RT) for de-protection of the linking functional groups, at a ratio of 1:1 plasmid to AuNP. Coupling of the plasmid to the AuNP surface was carried out by addition of the de-protected
plasmid to a solution containing the AuNP and allowed to place exchange for 48 h. The assembled AuNP-TSX/TSS/TSN-plasmids were pelleted out of solution by ethanol precipitation followed by centrifugation at 3000 rpm to remove unbound plasmid. Gel electrophoresis (1% agarose) of the AuNP-plasmid construct confirmed assembly and showed < 1% free AuNP present.

2.2.4 Transfection Agent

The AuNP transfection construct was prepared by encapsulating AuNP-SX-DNA, -SS-DNA, -SN-DNA, -TSX-plasmid, -TSS-plasmid, -TSN-plasmid into Lipofectamine2000® (Life Technologies). Lipofectamine2000® encapsulation of the AuNP constructs (AuNP was carried out at a concentration of 3 pmol of AuNP, as determined by the AuNP absorption plasmon at 525nm, to 10 µL of Lipofectamine2000®. The loading level of constructs into the liposome was chosen to be consistent with concentrations utilized in the protocol for a standard plasmid transfection in Lipofectamine2000®. Dynamic Light Scattering (DLS) data (DynaPro-Titan) was obtained at 20% laser power and used to analyze the size of the transfection packages. The hydrodynamic radius is calculated by averaging six replicates each containing 10 measurements with an acquisition time of 1 s, with the complexes suspended in nanopure water.

2.2.5 Cell Transfection

A375 human melanoma cells, cultured at 37°C with 5% CO₂ in Dubelco’s Modified Eagle’s Medium (DMEM-7777) (Sigma) supplemented with addition of 10% fetal bovine serum (Sigma) were plated at 30,000/cm² in 0.33cm² wells of a glass bottom 96-well plate or with 1.5 coverglass treated with poly-L-lysine (MatTek Corp). Cellular transfection was carried out 24 h after plating. Media was exchanged after 48 h of transfection to remove any unreacted AuNP complex. Cell viability was verified using Trypan blue exclusion test of cell viability after 24 h, in addition to visual conformation on the microscope to assess the cell health. No evidence of chromosomal condensation or changes in cell morphology was observed following transfection, indicative of no cytotoxicity at the transfection agent concentrations employed in this study. Trypan blue assays confirmed cell viability was maintained at 90% following transfection for all experiments relative to control cells. Cell division was occurring over the time span of the experiment, with a doubling rate of 2% per hour resulting in confluence at 48 h. Due to over-
confluence affecting cell growth behavior, the experimental data was limited to 48h for the uptake and release study and 72 h for the gene expression measurements. Experiments were conducted in triplicate.

2.2.6 Optical Microscopy

Confocal microscopy images were obtained on a Nikon Eclipse Ti Inverted Microscope at 20X magnification. Samples were excited using a Yokogawa automated 5000 rpm Spinning Disk Confocal equipped with lasers to excite the DyLight700 (640 nm, 900 ms exposure), DyLight488 (488 nm, 500 ms exposure), and tdTomato (561 nm, 400 ms exposure). Images were collected by an Andor Clara high resolution CCD at 20X magnification. Measurements were taken at 30 min intervals for 48 h while incubated in a stage mounted live-cell chamber, kept in focus by a Nikon Perfect Focus system. An additional measurement for gene expression was taken at 72 h post transfection. Images were analyzed for total intensity and cell number by ImageJ software. Intracellular uptake and release kinetics were monitored by tracking DyLight700 and DyLight488 intensities, respectively, normalized to cell count. Gene expression was assessed by the red fluorescent protein emission intensity from tdTomato.

2.2.7 Image Analysis

The kinetics of the images were analyzed with ImageJ software, using a standard rolling ball radius of 50 pixels to subtract the background to remove any cellular autofluorescence and the total intensity of each image was measured per time frame. The resulting intensity was corrected for cell count and plotted to an exponential one-phase model, to fit the first order association kinetics of the release of the sdDNA and plasmid from the surface of the AuNP, plotted with standard deviation of the samples. Each experimental trial was imaged in three separate regions per well in a 96-well plate per time point, with n = 3 per sample. The plots were analyzed for differences using the student’s unpaired t-test, to determine statistical differences in the samples. Protein expression was analyzed with ImageJ and measured fluorescence was corrected to cell count per time fame. Colocalization of the DL700 and DL488 was analyzed using the JACoP plugin for ImageJ to assess the van Steensel Cross Correlation Function (CCF) for red versus green channels, with best gaussian fit per sample calculated for the 12 and 24 h time points.
2.3 Results and Discussion

2.3.1 AuNP-DNA Constructs

The McSET-BOT incorporates two dye labels on a single duplex DNA to allow the cellular uptake (DL700) and subsequent intracellular disassembly (DL488) of the nucleic acid cargo appended to the 6 nm AuNP to be visualized. In Figure 2.1, the design of the McSET-BOT is shown for the 39 bp duplex DNA sequences, with the green DL488 “release sensor” appended to the DNA via a modified thymine (T) base at the 7th position from the AuNP surface; as well as the red DL700 “uptake sensor” appended at the 30th bp (Figure 2.1).

![Diagram](image)

**Figure 2.1** Scheme of AuNP-DNA coupling strategy, showing mono-thiol (SX), dual thiol (SS) and thiol plus amine (SN) with corresponding anticipated release times: t₁, t₂, and t₃, respectively. Cellular uptake is monitored by constant Dylight (DL) 700 emission, release from is monitored by turn on of DL488, according to SET theory, and gene expression is observed by emission of the fluorescent protein, tdTomato.

Although not explicit in Figure 2.1, ligation of a 6 kbp linearized plasmid, containing the gene for the fluorescent protein (FP) tdTomato, to the labeled 39-mer DNA duplex McSET-BOT probe allows for the reporting of uptake, timing of release, and correlation with gene expression...
rates. For the McSET-BOT, the DL700 provides a constant fluorescent marker of the AuNP cargo’s location, since it is not proximally quenched by the 6 nm diameter AuNP, and thus, is emissive even when bound to the AuNP surface. The DL488 operates as a fluorescence “turn-on” beacon to report the timing of intracellular release for the nucleic acid cargo from the AuNP surface. When the DNA is appended to the AuNP surface, the contact distance is within the >90% non-radiative energy coupling distance. The “turn-on” signal occurs at the moment of release due to a loss of the SET non-radiative energy coupling between the 6 nm AuNP surface and the excited state of DL488, once the DL488 is no longer bound to the AuNP and begins to diffuse away from the AuNP surface. Thus, as intracellular processing of the McSET-BOT results in nano-cargo release, and “turn-on” of the fluorescent marker allows for the kinetics of intracellular uptake and processing to be analyzed for each coupling strategy in real time, via live cell fluorescence microscopy.

The AuNP is prepared by the citrate reduction method. The citrate passivation layer is exchanged for bis-(sulfonatophenyl)phenyl phosphine (bSPP) prior to coordination of the functionalized nucleic acid sequences to aid in NP stability, by way of the more stable binding phosphine compared to carboxylates of citric acid. The bSPP also facilitates better DNA surface packing on the particle by providing a bulky tri-phenyl spacer molecule to prevent the DNA backbone from interacting with the surface of the AuNP, which would cause the DNA to lay flat rather than project perpendicularly to the AuNP surface, as is necessary for properly predicting SET behavior. TEM and size distributions of the as-synthesized AuNPs are shown in Figure 2.2A. The functionalized C₆-thiol (SH) and C₆-amine (NH₂) modified DNA sequences were synthesized commercially with DL488 and DL700 internal base labels on separate strands, with a 4 bp overhang to allow for ligation to the linearized plasmid containing the tdTomato gene and a cytomegalovirus (CMV) promoter to drive overexpression of the tandem dimer RFP within the cell. The McSET-BOT DNA sequence was duplexed by slow-cooled annealing procedures to produce nucleic acid sequences with monothiol (SX), dual thiol (SS), and thiol/amine (SN) binding moieties. For ease of reference, the McSET-BOT ligated to the tdTomato plasmid is abbreviated as TSX, TSS, and TSN, respectively.

In order to append the DNA to the AuNP, reduction of the synthetic dithiol protecting groups was accomplished by incubation in 5 mM tris(2-carboxyethyl)phosphine (TCEP), after which the synthetic DNA was added directly to the AuNP at a 20:1 DNA to AuNP ratio, with a
24-48h mixing period to allow for DNA attachment. The successful coupling of the DNA to the AuNP was evaluated by 1% agarose gel electrophoresis, which demonstrated retention of the AuNP-DNA complexes compared to the AuNP-bSPP due to the increased molecular weight of the attached DNA (Figure 2.2B).

**Figure 2.2** Characterization of the McSET-BOT AuNP showing (A) TEM (scale bar = 100 nm) and size distribution (inset) of the 6 nm diameter AuNP, prior to biofunctionalization; and (B) agarose gel electrophoresis of the as-synthesized AuNP (lane 1), monothiol DNA appended AuNP (lane 2), bidentate thiol appended DNA AuNP (lane 3), and bidentate thiol plus amine DNA appended AuNP (lane 4).

**Figure 2.3** Dynamic light scattering intensity as a measure of the hydrodynamic diameters for (A) AuNP-bSPP of ~14.2 nm, (B) AuNP-sdDNA of ~36.4 nm, and (C) AuNP-plasmid of ~166.7
nm. Mean values are shown as green filled bars and size distributions are shown in red unfilled bars.

Additionally, dynamic light scattering (DLS) was used to determine the change in hydrodynamic radii for AuNP-bSPP versus AuNP-sdDNA and AuNP-plasmid (Figure 2.3), showing an increase in size from AuNP-bSPP< AuNP-sdDNA< AuNP-plasmid, as is expected for successfully assembled constructs. Measured hydrodynamic diameters were found to be 14.2 nm, 36.4 nm, and 166.7 nm, respectively (Figure 2.3).

![Graph](image-url)

**Figure 2.4** Cyanide etching of the McSET-BOT. Dissolution of the AuNP is observed by loss of the 525 nm plasmon in the absorbance spectra (A) and release the DL488 and DL700 dye-labelled DNA is evidenced by the recovery in the fluorescence spectra (B) of DL488 at 520 nm. Absorbance of the dyes is shown at a 50x magnification of the absorbance intensity (inset). The DL700 emission at 700 nm is relatively unaffected due to non-SET behavior (C).
Verification of SET activity for the BOT constructs was performed by exposing assembled complexes to cyanide etching with a 0.1 M NaCN solution which was able to dissolve the AuNP, liberating the bound DNA and allowing for fluorescence recovery of the attached fluorophores. Dissolution of the AuNP can be seen by the loss of the AuNP absorption plasmon at 525nm (Figure 2.4A), which releases the attached DNA to recover fluorescence of the quenched DL488 (Figure 2.4B). The DL700 signal was unaffected, demonstrating the unquenched non-SET relationship between the DL700 and the 6 nm AuNP, as expected since the DL700 emission is lower in energy than the AuNP plasmon and thus non-energetically coupled at the distance it is placed from the AuNP surface on the attached DNA strand (Figure 2.4C).

2.3.2 Cell Transfection and Imaging

The impact of the surface binding functionality (SX, SS, SN) on the dynamics of intracellular McSET-BOT nanoparticle uptake and nucleic acid release was analyzed in Human melanoma cells, A375. The A375 cells were plated in 96-well plates with glass bottoms for optimum fluorescence light collection. A375 cells were transfected 24 h after plating with AuNP-SX/SS/SN-DNA and AuNP-TSX/TSS/TSN-plasmid complexes using Lipofectamine2000® transfecting reagent, following media exchange. Cells were incubated at 37°C using a live cell chamber mounted on the microscope stage, and live cell fluorescence images were collected every 30 min for 48 h. From the selected time points displayed in Figure 2.5, it can be seen that the DL700 fluorescence signal is continuous throughout the experiment, demonstrating no SET quenching and allowing for visualization of the location of the AuNP-DNA throughout the experiment. Conversely, the DL488 fluorescence is “off” at early time points and gains brightness over time as more DNA is released from the surface of the AuNPs, making apparent the slight differences in the time evolution of release for each construct. It is observed in Figure 2.5 that at early time points, fluorescent packages are seen with punctate patterning inside of the cells, indicating endosomal packaging, as is expected with the use of the cationic liposome Lipofectamine2000® transfection reagent. At later time points, more cells display diffuse fluorescence intensity throughout, indicating endosomal escape of the dye-labeled DNA and diffusion into the cytosol. Inspection of the intensity patterns in Figure 2.5 reveal the order of release follows the expected behavior for binding affinity SX>SN>SS, reflecting the number of chelates to the AuNP surface and the binding strength of an amine relative to a thiol.
Figure 2.5 Live cell fluorescence microscopy of A375 cells incubated with (A) AuNP-SX, (B) AuNP-SS, and (C) AuNP-SN over 48 h. DL700 denotes uptake and particle tracking, DL488 shows DNA release, DIC shows normal cell morphology, and overlay shows localization. Scale bar: 50 µm.
Figure 2.5 - continued
Figure 2.5 - continued
2.3.3 Cellular Colocalization Analysis of DL488 and DL700

To assess the stability of the McSET-BOT constructions and ensure no degradation of nucleic acids occurred during the experimental time frame, analysis of the colocalization of the DL488 and DL700 fluorescent labels was conducted. Assessment of the cross correlation function (CCF),149 as described by van Steensel and coworkers, provides the most unbiased measure of colocalization for fluorophores with punctate fluorescence patterns, especially when the fluorophores possess differential fluorescence quantum efficiencies and detector sensitivities for their emission windows.140 In CCF the Pearson’s correlation coefficient of colocalization ($r_p$, Eqn. 2.1) is calculated for and compared between two potentially colocalizing fluorescence channels, “A” (DL700), versus “B” (DL488), as they are laterally shifted by 20 pixel increments.

$$r_p = \frac{\sum (A_i-a) \times (B_i-b)}{\sqrt{\sum (A_i-a)^2 \times \sum (B_i-b)^2}}$$

(Equation 2.1)

Where $i$ denotes intensity values for a given pixel, and a & b denote the intensity averages over the whole image per channel, A and B.151 The best Gaussian fit of the plot created by this lateral pixel shifting of the fluorescence images versus the calculated effect on the Pearson’s correlation coefficient gives the CCF.149,152 Conclusions about the colocalization are drawn from the overall shape and amplitude of the CCF plot, where maxima at the 0 pixel shift position indicates the presence of colocalization, and the amplitude (0-1) indicates the degree of spectral intensity matching between the channels.149,152

CCF analysis of the DL488 and DL700 fluorescence channels for each McSET-BOT construct is shown in Figure 2.6 at 12 h and 24 h after transfection, for the short DNA and plasmid genes attached to the AuNP by the various coupling strategies. It can be seen that all the coupling strategies for both short and full plasmid DNA show remarkable colocalization at the 12 h and 24 h time points (Figure 2.6A-F and G-L), with strong Gaussian distributions for the CCF of all samples, indicating no separation of fluorophores due to nuclease cleavage of the DNA strands.

2.3.4 Uptake and Release Kinetics

In Figure 2.7, a plot of the time dependent change in red and green fluorescence intensity is shown. The change in intensity is a measure of the kinetics of uptake (via DL700 intensity)
Figure 2.6 Analysis of colocalization for DL488 and DL700 fluorescence, expressed in Cross correlation functions (CCF) for AuNP-SX/SS/SN-DNA (A, C, E and G, I, K) and AuNP-TSX/TSS/TSN-Genes (B, D, F and H, J, L) at 12 h and 24 h post transfection, respectively. Best Gaussian fit is shown in blue.
Figure 2.6 - continued
and release (via DL488 intensity) for the labelled DNA. Images were processed in FIJI ImageJ software to collect total fluorescence for each channel, and the kinetics were calculated in Igor software, according to the sigmoidal plot function. The calculated intracellular release kinetics for the AuNP-SX/SS/SN-DNA and AuNP-TSX/TSS/TSN-plasmid coupling strategies are displayed below (Table 2.1).

**Figure 2.7** Kinetic data of AuNP uptake (A & C) and cargo release (B & D) for the short duplex DNA (open shapes) and plasmid gene (filled shapes) coupled to the AuNP by monothiol “SX” (black circle), dual thiol “SS” (red triangle) and thiol plus amine “SN” (blue square), respectively. Uptake is monitored by cellular fluorescence of DL700 and Release by DL488, normalized to cell counts. Trend lines reflect sigmoidal fits.
The timing of release for each DNA sequence follows SX>SN>SS for the speed of intracellular disassembly. Changes in uptake patterns between the different assembly strategies, while unanticipated, may reflect differences in overall DNA loading onto the AuNP surface and related charge density affects within the cationic liposome formation, creating differentially sized liposomes which may experience non-uniform rates of cellular internalization due slower diffusion of larger liposome towards the cellular surface.

### Table 2.1 Intracellular DNA cargo release kinetics from 6nm AuNP for monothiol (AuNP-SX-DNA and AuNP-TSX-plasmid), dual thiol (AuNP-SS-DNA and AuNP-TSS-plasmid), and thiol plus amine (AuNP-SN-DNA and AuNP-TSN-plasmid) attachment, for short DNA oligonucleotides and linearized plasmids, respectively.

<table>
<thead>
<tr>
<th>AuNP-Construct</th>
<th>Bond half-life (h)</th>
<th>Release Rate (%•h⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SX-sdDNA</td>
<td>7.9 ± 4.9</td>
<td>6.33</td>
</tr>
<tr>
<td>SS-sdDNA</td>
<td>15.9 ± 1.5</td>
<td>3.15</td>
</tr>
<tr>
<td>SN-sdDNA</td>
<td>12.2 ± 2.0</td>
<td>4.10</td>
</tr>
<tr>
<td>TSX-plasmid</td>
<td>8.0 ± 4.9</td>
<td>6.25</td>
</tr>
<tr>
<td>TSS-plasmid</td>
<td>15.4 ± 1.5</td>
<td>3.25</td>
</tr>
<tr>
<td>TSN-plasmid</td>
<td>10.6 ± 1.9</td>
<td>4.72</td>
</tr>
</tbody>
</table>

Interestingly, in the A375 cells the release dynamics for short DNA *versus* the plasmid is nearly identical. Based on earlier studies, it was expected that the timing of release would vary between the DNA lengths due to the added molecular weight and charge density of the large plasmid compared to the short DNA oligonucleotide. This was not observed, most likely due to a reduction of the DNA loading levels in these studies.

### 2.3.4 Coupling Influence of Timing of Fluorescent Protein Gene Expression

To analyze the surface coupling dependent release and the effect on the timing and extent of gene expression of the TSX, TSS, and TSN linearized plasmid bound onto the AuNP was compared to lipofectamine delivered circular plasmid and the TSS linearized plasmid in the absence of a AuNP (Figure 2.8). The nano-delivered plasmid carries the gene for the fluorescent protein tdTomato as a model of a therapeutic gene. Specifically, tdTomato was chosen for its brightness, as it is a tandem dimer of the mCherry fluorescent protein and thus twice as luminous, making it highly detectable via fluorescence microscopy.
Figure 2.8 (A) Live cell fluorescent microscopy images of the tdTomato fluorescent protein gene expression from 24-72 h post transfection, as delivered by the AuNP via mono thiol (TSX), dual thiol (TSS) or thiol plus amine (TSN) DNA linkage. Scale bar is 50 μm. (B) Quantification of cellular gene expression from 24-72 h post transfection for the AuNP-TSX, AuNP-TSS, and AuNP-TSN attachment strategies, along with the circular plasmid and a linearized plasmid functionalized with dithiol protected dual thiol modifications, as controls.
Expression of the tdTomato gene is signaled by whole-cell red turn-on, as the protein is cytosolically overexpressed by the CMV promotor, built into the plasmid backbone. In Figure 2.8A, the progression of fluorescent protein expression can be seen for the various AuNP delivery constructs from 24 to 72 h. From the microscope images, TSX is observed to maintain a somewhat constant expression level after 36 h, whereas both TSS and TSN exhibit a steady increase in tdTomato brightness in time.

The delay in expression is longest for the TSS sequence consistent with the delayed intracellular release observed in Figure 2.5, and consistent with the expected binding strength of the dual thiol linkage chemistry. Quantification of the expression rates versus circular plasmid and linearized plasmid (TSS) further emphasizes the differences in expression rate for the AuNP-TSX, -TSS, and -TSN, where for AuNP-TSS the level of expression is significantly greater at 72 h than observed at 36 h, in contrast to the single thiol and thiol plus amine, as well as the circular or linearized plasmids without AuNPs (Figure 2.8B).

2.4 Conclusion

This study demonstrates that by utilizing differential surface appendage strategies for coupling therapeutic cargo, such as short DNA oligonucleotides and linearized plasmids, to 6 nm AuNPs, distinct release kinetics can be achieved for thiol versus amine and monodentate versus bidentate. It was found that the monothiol linkage to AuNP releases fastest, followed by thiol plus amine attachment, with dual thiol attachment being the most stable and taking the longest to release. This is believed to be due to the availability of π-orbitals which can engage in back bonding with the surface of the AuNP. Interestingly, mono amine and dual amine attachment strategies were not found to be chemically stable for DNA attachment to the AuNP and readily dissociated during clean up and characterization (data not shown).

It was also found that the length of the attached DNA did not factor into the release rates for the cell line observed, where the large molecular weight (6 kbp) plasmid did not take longer to release in comparison to the short (35 bp) oligonucleotide, as was expected, but rather the general trend of release was found to be: SX/TSX>SS/TSS>SN/TSN, which was consistent for both sizes of DNA cargo. Understanding and utilizing AuNP surface chemistry that is able to produce controllable conjugation of therapeutics to the nano-medicine allows for tunable release of medicine in the
cellular environment, getting one step closer to personalized medicine. Ideally, the modulation of nano-cargo release for gene therapy applications would allow for a greater than 12 h time differential between one cargo release event and the next, to give ample time for the first gene in the therapeutic sequence to release, express, and build up sufficient intracellular concentration prior to the second therapeutic agent’s release (~12 to 24 h delayed). Possessing these tools will open the possibility for coordinated therapies, such as in the treatment of multi-drug resistant cancers. For example, a gene for efflux pump knock-down could be delivered, followed by an apoptotic gene and chemotherapeutic drug delivery, in order to disable the cancer cell’s native defense mechanisms and specifically destroy the cancer cells, with no non-target side effect, as are so common with modern pharmaceutical approaches.
CHAPTER 3

A GOLD NANOPARTICLE BIO-OPTICAL TRANSPONDER TO ACTIVELY
MONITOR INTRACELLULAR pH


3.1 Introduction

The interfacing of gold nanoparticles (AuNP) with surface appended biological polymers (DNA, RNA, protein, antibodies) has led to the design of probes capable of reporting subtle structural perturbations in biological systems,

54,58,131,154,155 detecting analytes in complex media.57,156–162 Most notably, bioconjugated AuNPs have been employed as bio-optical transponders (BOTs) that report the intracellular level of reactive oxygen species (ROS) in endosomes,163 the timing of therapeutic nucleic acid (NA) payload release following transfection,132 changes in metabolite levels within cells,156,160,164 and to even allow dynamic monitoring of coordinated cellular responses.74,101,102,132,165–170 While the gold nanoparticle BOT technology has developed rapidly, the observations that therapeutic nucleic acid sequences (siRNA, antisense DNA, genes) delivered on a gold BOT may disrupt endosomal maturation, exhibit resistance to enzymatic degradation, and experience enhanced activity, are not well understood.100,171,172 Exploring the intracellular processing and fate of nano-delivery agents is a necessity for intelligent nanomedicine development, as the nano-agent must be able to successfully and effectively deliver its therapeutic cargo to the intended intracellular target.

In this chapter, a live-cell pH-BOT is developed, capable of dynamically reporting the evolving intracellular pH of the endo-lysosomal pathway following transfection with a duplex DNA functionalized 6.6 nm AuNP, enabling temporal resolution that is unachievable in fixed cell samples. The pH-BOT sensor reports the NP uptake, DNA cargo release, and the intracellular pH experienced by the cargo, using automatically collected live-cell confocal fluorescence microscopy imaging (Figure 3.1). The pH-BOT sensor operates by utilizing a three-dye Multi-color Surface Energy Transfer (McSET)53,58,130,173 probe design which provides a barcoded optical response from three spectrally distinct fluorophores. NP uptake is reported by DyLight 700 (DL700) a non-SET dye, while DNA cargo release following endosomal packaging and cellular processing is reported by the fluorescence turn-on of a pair of SET-active dyes,
Figure 3.1 Schematic representations of (A) the endo-lysosomal pathway for nanoparticle cellular uptake, showing thermal plot of pH processing, starting with endocytosis and transport inside a trafficking vesicle at near neutral pH (red) to (i) the Early Endosomal (EE) at a pH between 7-6 (red-orange), before processing to (ii & iii) a Late Endosome (LE) acidifying from pH 6-5 (yellow-green), and finally trafficking to (iv) the Lysosome (LY) for acidic degradation at pH 5-4.5 (blue-purple), with potential exocytosis of recycled materials at pH 7 and above (red). And (B) the 3-color pH-BOT probe, displaying 6.6 nm diameter gold nanoparticle (AuNP), with appended thiol-modified 45mer oligonucleotide possessing dye-labels of an unquenched DL700 at the 45th bp (uptake sensor), a SET quenched & weakly pH-sensitive DL405 at the 32nd bp (release sensor), and a SET quenched & highly pH-sensitive FAM at the 11th bp (pH sensor).
fluorescein amidite (FAM) and DyLight 405 (DL405). The pH-BOT probe design is based on surface energy transfer (SET) methods to access the advantages of using a gold nanoparticle as an energy acceptor and to simulate gold nanoparticle therapeutic delivery into cells. SET based optical ruler methods are enhanced over traditional Förster’s Resonance Energy Transfer (FRET) techniques, whether molecular or quantum dot based, due to the efficiency of quenching of dyes\(^\text{26}\), which greatly reduce the rates of laser-induced dye bleaching in multi-hour live-cell experiments. Additionally, the greater distance of energy coupling achievable in SET allows for dye-labelled nucleic acid molecules to be utilized as therapeutic mimics to more accurately investigate the processing of nanotherapeutic delivery systems. And functionally, the large optical cross section for a 6.6nm AuNP (~10\(^5\) for a fluorophore, as compared to ~10\(^9\) for 6 nm diameter AuNPs), coupled to the sharp drop in absorption strength for AuNPs at wavelengths longer the plasmon peak (i.e. fluorescence quenching does not affect fluorophores red of the AuNP plasmon), allow the AuNP to possess band-pass like behavior. The combination of these features is essential to the function of the pH-BOT and allows the on/off quenching of the broadly emitting pH-cassette (DL405-FAM), while permitting the unmodified emission of the tracer dye (DL700). SET calculations determined that synthesized 6.6 nm diameter gold nanoparticles fall within the ideal NP size range that would completely quench the pH responsive dye cassette (consisting of FAM at bp 11, and DL405 at bp 32 relative to the particle surface), while leaving the tracer emission unquenched (DL700 at bp 45).

The rapid development of therapeutic methods utilizing a plethora of nanoparticle delivery vehicles has resulted in the appearance of several pH based nanosensors, or pH-triggered nano-delivery agents, incorporating FRET based dye labels onto mesoporous silica and graphene nanoparticles,\(^\text{109,174-177}\) squarine dyes on CdSe@ZnS quantum dots,\(^\text{178,179}\) dye labelled hyperbranched polymers,\(^\text{180-182}\) upconversion nanoparticles,\(^\text{183,184}\) and nanoparticle-aptamer systems.\(^\text{115,185}\) These breakthrough studies established the ability to use and identify pH via ratiometric measurements and recently have been applied to cell environments, although no dynamic live cell imaging coupled to monitoring of therapeutic delivery of a nucleic acid via a gold nanoparticle has been demonstrated. The development of a AuNP-based pH sensor capable of correlating the delivery and processing of nucleic acid agents entering a cell will answer questions pertaining to the fate and transport of nucleic acid therapeutic, once transfected into a cell. Creation of this environmentally responsive nucleic acid therapeutic probe requires the
reporter dye labels to be co-assembled onto NA sequences that mimic therapeutic nucleic acids models, such as siRNA delivery. Such studies are highly relevant in response to the rise of gold nanoparticle based anti-cancer nanotherapeutics already advanced through Phase I clinical trials (Aurimune®).

Utilizing insights gleaned from explorations in the field of cellular biology, particularly endo-lysosomal maturation and acidification progression, an investigation of the effects AuNP therapeutic transfection and cellular processing versus the normal cell cycle is possible. The SET based gold nanoparticle pH-BOT provides a dynamic assessment of the evolution of endosomal pH as the endosome matures from early to late, and then to a lysosome, by reporting the ratiometric intensity of FAM to DL405. The experimental results support a model where the nanotherapeutic is endosomally encapsulated in <1 h of transfection, with release of the DNA duplex cargo with a half-life of 1.5 h. As the endosome matures, acidification occurs but there is an observed asymptote at pH 6.0 for up to 24 h, suggesting a stalling at the late endosome stage. Image analysis of stained early and late endosomes, and lysosome packages reveal that although the pH appears to be buffered, colocalization of the pH-BOT is observed in all components of the endo-lysosomal pathway. It is believed the asymptotic behavior reflects a pH buffering effect attributed to the protonation of the alkyl thiolates (pKa > 10) used to couple the DNA to the gold surface once released following endosomal packaging. The effect of pH buffering may explain the earlier reports of enhanced functionality of the nucleic acid cargoes and enhanced stability due to reduction of NA degradation pathways. The study provides further insights into endosomal maturation and the cellular processing of nanotherapeutics. While maturation of endosomes carrying pH-BOT is slowed significantly, finding that pH-BOT is distributed throughout the endo-lysosomal system while remaining at an average pH of ~6 indicates that a slow, but persistent maturation continues to occur. This observed decoupling of endosomal maturation from acidification lends support to those models which propose that pH alone is not sufficient to explain endosomal maturation and the progressive exchange of Rab proteins.

The developed pH-BOT will open doors for diverse cell delivery inquires and generate greater insight into our understanding of the fundamental processes of biology. Being able to monitor the pH evolution will allow for design of nanotherapeutic agents to ensure escape of the cargo from endosomes before full acidification occurs which is known to lead to NA damage
through depurination of NAs at low pH. The results of the study indicate the pH-BOT can actively monitor the pH during delivery of NA therapeutic cargos. The pH-BOT incorporates dye labelled-dsDNA on an AuNP as a mimic of therapeutic AuNP@NA delivery systems common in biomedical research and similar to other reported NP@NA formulations. By attaching the reporter fluorophores directly to the delivered NA the progress of NA cargo can be tracked through the endo-lysosomal system, allowing for a more comparable study of NA delivery from the AuNP than afforded by similar efforts in the literature focused on tracking nanotherapeutic delivery of small-molecules and endosomal sorting after uptake. The pH-BOT models NA delivery and fate explicitly, while providing reporting of cellular uptake and the moment of release from the AuNP, as well as monitoring the location of the NA cargo within the cells not only spatially but also in identifying the containing endosome by the stage of endosomal maturation, to better elucidate NA cellular processing when delivered from AuNPs. In contrast to other studies using pH-sensitive dye-cassettes to monitor endo-lysosomal maturation, the pH-BOT uses SET to generate a ratiometric pH cassette that only turns ‘on’ after AuNP release; allowing for the tracking the endosomal state beginning at the moment of cargo release, followed by sorting and escape into the cytosol. Additional inclusion of the ‘always-on’ dye, DL700, which does not experience AuNP quenching from a SET interaction, enables tracking of uptake, sorting, and NA integrity; while simultaneously providing a means to measure DNA cargo release rates ratiometrically.

3.2 Materials and Methods

All chemicals were purchased through MilliporeSigma, unless otherwise noted.

3.2.1 AuNP Synthesis

Spherical 6.6 nm diameter AuNPs were synthesized by the citrate method according to published protocols. Briefly, a sparged aqueous solution of tetrachloroauric acid hydrate (Strem Chemicals) was reduced in the presence of citric acid and tannic acid at 60 °C with rapid stirring at a ratio of 1:4:0.5 mL of each 1% (by mass) solution in 100 mL of H2O, total volume. The prepared AuNPs were exchanged with bis(p-sulfonatophenyl)phenylphosphine (BSPP) as the protecting group. Particle size and shape analysis was conducted using a JEM-ARM200cF Transmission Electron Microscope for imaging and ImageJ for manual measurement of the
diameters of 100 particles. Size distribution and Gaussian fit of 6.6±1.0 nm were calculated in Igor software. DNA biofunctionalization of the AuNP was conducted by stable displacement of the coordinated BSPP passivation layer through thiol place exchange reactions using thiol-terminated DNA sequences. The synthetic DNA probe sequences are given below.

3.2.2 DNA Probes

Oligonucleotide sequences were designed to avoid FRET interactions of the dyes with one another, as well as produce optimal SET interactions for specific turn on of the release and pH probes and continuous fluorescence (non-SET coupling) for the uptake probe. The oligonucleotides were purchased from Midland Oligos (Midland, TX), with the sequences given below (modified bases are underlined).

Uptake probe strand: 5’-(thiol-C6)CGA GGT TGG CAA GAG TAA CAC GGA ATT CAG TAT GCA CAC GGA ACG(DyLight 700)-3’

pH/release probe strand: 5’-CGT TCC GTG TGC A(DyLight405)TA CTG AAT TCC GTG TTA CTC T(FAM)TG CCA ACC TCG-3’

A calibration curve was created to measure the ratiometric absorbance and fluorescence response of the dye-labeled DNA sensor to changes in pH from 4-9 using McIlvaine’s buffer, via a Cary Bio 50 UV-Vis spectrophotometer (data not shown) and a Cary Eclipse fluorometer.

3.2.3 Synthesis and Characterization of the 3-color pH-BOT Probe

As previously described, the thiolated duplexed DNA probe sequence was appended to the AuNP via thiol place exchange with the aid of 5 mM tris(2-carboxyethyl) phosphine (TCEP) to reduce the protected dithiol on the synthesized DNA, followed by 24 h reaction of the DNA and AuNP in a 20:1 ratio at RT. After which the complex was isolated by centrifugation in ethanol and resuspended in Nanopure™ H₂O. The complex was then characterized by 1% agarose gel electrophoresis. The amount of dye-labeled DNA was quantified by UV-Vis absorption spectroscopy and the amount of SET quenching was quantified by cyanide etching of the AuNP to determine fluorescence recovery.
3.2.4 Investigation of pH-BOT Probe Buffering and Release in Cuvette

The pH-BOT probe was incubated at RT or 37°C in McIlvane’s buffer solutions ranging from pH 3.9 to 7.4 and CHO cell lysate solutions at pH 5 and 7, and fluorescence response was measured using a Cary Eclipse spectrophotofluorometer over 24 h. CHO cell lysate was prepared by three-fold rapid freeze-thaw lysis of ~1 million CHO cells in phenol red-free culture medium, followed by addition of 0.5 M EDTA to impair nuclease activity. The cell lysate was then subjected to a 0.2 μm syringe filtration to remove cell membrane aggregates, and pH adjusted with 1 M HCl or NaOH. Calculated pH values were obtained from the integrated fluorescence intensities ratios of FAM (excitation 480 nm) to DL405 (excitation 390 nm) using the calibration equation, computed in MATLAB (MathWorks®).

3.2.5 Cell Culture and Transfection

Chinese hamster ovary (CHO) cells acquired from ATCC were cultured at 37°C and 5% CO2 with 50% humidity were grown in Dubelco’s modified Eagle’s medium (DMEM 2902) supplemented with non-essential amino acids, 10% cosmic calf serum (CCS) and antibiotics (Gibco), and were plated at 30,000 cells/cm² in glass bottom 96-well plates (MatTek) 24 h prior to experimental transfection. McSET pH probe complexes were encapsulated in Lipofectamine2000® (Life Technologies) according to optimized manufacturer instructions, briefly 1 µL of Lipofectamine® reagent was used per 2 pmol of pH-BOT in a total volume of 60 µL of serum free FluoroBrite imaging media (Life Technologies), incubated together at room temperature for 20 mins immediately prior to transfection, of which 20 µL was added to fresh serum containing FluoroBrite imaging media per well. Endo-lysosomal vesicle tracking was conducted by treating cells with CellLight™ Early Endosomes-RFP, BacMam 2.0 (Molecular Probes), CellLight™ Late Endosomes-RFP, BacMam 2.0 (Molecular Probes), or LysoTracker DND-99 (Molecular Probes), prior to transfection with the pH-BOT probe according to manufacturer instructions.

Fixed cell slides were prepared at 2 and 6 h post transfection with CellMaskTM Orange membrane stain (Molecular Probes) to assess internalization of the pH-BOT, according to the manufacturers protocol.
3.2.6 Optical Microscopy

Confocal fluorescence images were collected using a Nikon Eclipse Ti inverted microscope at 40x and 20x magnification. Samples were excited using a Yokogawa automated 5000 RPM spinning disk confocal microscope equipped with lasers to excite the DL405 (405 nm, 900 ms exposure), FAM (488 nm, 500 ms exposure), and DL700 (640 nm, 900 ms exposure). Images were collected by an Andor Clara interline CCD high resolution camera. Measurements were taken at 30 min intervals for 24 h while incubated in a stage-mounted live cell chamber (kept at 37 °C, 5% CO₂ and 50% relative humidity). Focus was maintained throughout the time course of the experiment using a Nikon Perfect Focus system.

Membrane-stained fixed cells were imaged using a Zeiss LSM 880 inverted confocal microscope. Fluorophores were excited with 405 nm (DL405), 488 nm (FAM), 543 nm (CellMask™ Orange), and 633 nm (DL700) lasers, using a Plan-Apochromat 63x/1.40 Oil DIC M27 objective.

3.2.7 Image Analysis

Images were analyzed for fluorescence intensity and cell number by ImageJ and Nikon Elements software, with construct tracking of DL700 intensity and intra-endosomal pH at the time of release and during cell processing monitored by tracking the FAM to DyLight405 ratio of intensities, for punctate fluorescence regions of interest.

Automated image analysis was carried out in ImageJ software, utilizing the thresholding function to isolate endosomal fluorescence signals from the background intensities. Analysis of the intensity ratios of DL405 to DL700 gave rise to the rate of release from the NP surface over time. Analysis of the FAM to DL405 ratios showed the effect of changing endosomal pH on the pH-sensitive FAM over time. Experimental N of 5 replicates, with 3 imaging fields per replicate.

Analysis of the endosomal populations was performed using Nikon Elements Analyst software. Individual endosomes were selected using a region of interest tool, and ratios of FAM to DL405 were extracted for each individual endosome. Experimental N of 4 experimental fields, with analysis conducted on approximately 250-500 individual endosomes per time point.

Fluorescence intensity ratios were converted to pH according to the calibration equation (Eqn. 3.1) for the pH-BOT probe, using MatLab software. Histograms of endosomal pH
populations were calculated in Igor software, as were sigmoidal and linear fits for the calibration curve and DNA release kinetics.

3.2.8 Endo-Lysosomal Tracking Colocalization Analysis

Fluorescence images were deconvoluted to remove background fluorescence noise contamination with the rolling ball background subtraction of 50 pixels in ImageJ. Colocalization was assessed by comparison of the RFP channel for CellLight™ RFP-Rab or LysoTracker™ DND-99 for each vesicle with the DL700 channel for the pH-BOT probe via JACoP (ImageJ plugin)\textsuperscript{152} to assess the cross-correlation function and Manders’ colocalization coefficient for each image set. Thresholds were set for each image and channel individually. Averages and standard deviation were calculated in Microsoft Excel.\textsuperscript{TM}

3.3 Results and Discussion

3.3.1 Nanotherapeutics in the Endo-Lysosomal Pathway

Trafficking of therapeutic cargo delivered by a gold BOT into a cell typically occurs via the mechanism of endocytosis, resulting in rapid cellular packaging and processing through the endo-lysosomal pathway (Figure 3.1).\textsuperscript{207} The endo-lysosomal cellular defense response is marked by a rapid acidification of encapsulated foreign material through upregulation of V-ATPase proton pump activity as maturation of the endosome from early endosomes (EEs), to late endosomes (LEs), and finally lysosomes (LYs) occurs.\textsuperscript{85} The EE has a typical pH of approximately 7-6, as well as a membrane containing Rab5 proteins\textsuperscript{85,208} (Figure 3.1A i). As the endosome and its contents are processed by the cell, membrane fusion and fission events occur which allow for gradual acidification of the endosome and replacement of the Rab5 proteins for Rab7 proteins, which engender the more mature endosome to have the classification of LE, also referred to as a multi-vesicular body, with a pH of approximately 6-5 (Figure 3.1A ii & iii).\textsuperscript{209,210} LEs eventually acidify and fuse with LY which are capable of acidifying to a pH of 5-4.5 (Figure 3.1A iv). The maturation of the endosome represents one of the primary cellular defense mechanism against viruses and other pathogens, thus the maturation of the endosome leads to efficient nucleic acid degradation and thereby the loss of therapeutic potency for gene therapeutics.\textsuperscript{211}

For siRNA, antisense DNA, or other gene therapeutics to realize their full therapeutic
benefit, the intact nucleic acid sequences must depart the endo-lysosomal pathway prior to full acidification of the late endosome and subsequent fusion with the lysosome, where exogenous material is broken down into its rudimentary amino acid and nucleotide building blocks, if no previous escape or shunting transport has occurred.

3.3.2 Design of the pH-BOT Probe

To report the pH experienced by the nanotherapeutic agent once transfected into Chinese hamster ovary (CHO) cells, a live-cell bio-optical transponder was designed to actively report the pH (pH-BOT) following release of an AuNP delivered NA cargo within the endo-lysosomal package. The pH and nanotherapeutic location is monitored by a fluorescent dye cassette chosen specifically to provide spatial and temporal information by colocalization reporting. The pH-BOT is designed around a spherical AuNP, with a diameter of 6.6 nm, that has been biofunctionalized with ~20 DNA per AuNP (Figure 3.1B). The remaining sites are passivated by bis(p-sulfonatophenyl)phenylphosphine (BSPP). The 45 basepair (bp) duplex DNA is synthetically prepared to couple three dyes: DyLight (DL) 405 as an internal base modification of the thymine (T) at the 32nd bp, fluorescein amidite (FAM) as an internal T base modification at the 11th bp, and DL700 as a 5’ phosphate appendage at the 45th bp relative to the AuNP surface. The DL405 and FAM are on the same single strand of DNA, while the DL700 is bound to the complimentary strand, to monitor duplex denaturation and DNA degradation. The three dyes are coupled through C₆ spacers and are chosen to continuously monitor cellular processing and ensure colocalization analysis can be performed to eliminate errors from nuclease degradation of the pH-BOT. Transmission electron microscopy (TEM) images confirm the AuNP size of 6.6 nm with a spherical morphology (Figure 3.2 A & B). Confirmation of binding is provided by gel mobility studies using 1% agarose gel electrophoresis (Figure 3.2C). Indicative of DNA binding to the AuNP via the reduced thiol linkage, the gel shows the pH-BOT is retained relative to the as-synthesized AuNP, and AuNP in the presence of protected dithiol DNA (RS-S-DNA). Upon binding the tri-labelled duplex DNA to the AuNP surface, the emission from FAM and DL405 is observed to be quenched, while the emission intensity for DL700 is unchanged (Figure 3.3A). Further evidence of coupling is demonstrated by sodium cyanide etching of the AuNP to release the duplex DNA, resulting in immediate turn-on of the
FAM and DL405 emission signatures (Figure 3.3A), and dissolution of the AuNP can be seen by the loss of the plasmon at 525 nm in the absorbance spectra (Figure 3.2D).

**Figure 3.2** Nano-bioconjugate characterization showing (A) TEM of as-synthesized spherical AuNPs with (B) Gaussian fit to the size distribution of 6.6 ± 1.0 nm. (C) 1% agarose gel electrophoresis of the AuNP pH-BOT probe (i), versus as-synthesized AuNP with protected RS-S-DNA (ii), and as-synthesized AuNP with no DNA (iii). Retention of the AuNP pH-BOT probe versus controls demonstrates reduced thiol facilitated DNA binding to the AuNP. (D) Absorbance spectra of sodium cyanide etching of the AuNP away from the pH-BOT probe, with loss of the AuNP plasmon at 525 nm indicating cyanide dissolution of the AuNP and liberation of the previously bound duplex DNA strands, and (inset) 50x zoom of post cyanide absorption showing DL405, FAM and DL700. Absorption of AuNP displays a concentration of 0.01 µM ($\varepsilon_{525}=1.97 \times 10^7$ M$^{-1}$ cm$^{-1}$) prior to cyanide treatment, and the duplex pH-BOT DNA gives a concentration of 0.19 µM ($\varepsilon_{260}=7.3 \times 10^5$ M$^{-1}$ cm$^{-1}$) after cyanide.
3.3.3 Optical Response of the pH-BOT

The DL700, DL405 and FAM labels provide three functions: i) tracking cellular uptake of the pH-BOT, ii) spectrally reporting release of the DNA from the AuNP surface by a photoluminescence turn-on event due to loss of SET quenching as the DNA diffuses away from the AuNP surface, and iii) providing an active pH reporter by ratiometric analysis of FAM to DL405 fluorescence intensities, reflecting the pH sensitivity of FAM and near insensitivity of DL405 for the physiologically relevant pH range of the endo-lysosomal pathway (pH 7.5 to 4.5). FAM is highly pH sensitive due to its three protonatable oxygens providing pKa points at 2.2, 4.3, and 6.7. In comparison, DL405 is weakly sensitive to pH, making it a good ratiometer for measuring intracellular pH (Figure 3.3B). Optical tracking of the pH-BOT from uptake to endo-lysosomal processing is achieved by monitoring the emission of DL700. DL700 photoluminescence is not quenched in the presence of the AuNP allowing its intensity to provide a continuous monitor of the cellular location and packaging of the pH-BOT. Since DL700 is relatively pH insensitive over the physiologically relevant range (Figure 3.3B), it provides an internal intensity standard for analysis of the integrity and concentration of active pH-BOT. Observation of colocalization of DL700, with DL405 and FAM emission, as well as constant intensity of DL700 during the experimental measurements, signal the absence of nuclease induced DNA degradation or acidic depurination of the DNA. As observed in cuvette studies, the ratio of DL405 to DL700 remains relatively unchanged between pH 5-7.5, and can be used as a ratiometric measure of intracellular DNA release (Figure 3.3D).

The release of the DNA cargo from the AuNP surface and pH of the environment experienced by the cargo is reported by the emission profile of FAM and DL405. DL405 and FAM are quenched when bound to the gold surface through surface energy transfer (SET) mechanisms. Following the scission of the Au-thiolate bond as the endosome acidifies, the release of the duplex DNA into the endosomal lumen is reported by the ratiometric turn-on of emission for FAM to DL405. In Figure 3.3A, the optical response of the DNA release event is calibrated in cuvette studies wherein the dye-labeled DNA is liberated from the AuNP by cyanide dissolution causing fluorescence recovery of the quenched FAM and DL405.

The pH environment of the pH transponder DNA duplex is reported ratiometrically by the intensity ratio of FAM relative to DL405. A calibration curve was generated by plotting the response of the pH-responsive DNA sequence in the absence of the AuNP in various pH buffers.
Figure 3.3 (A) Emission spectra and schematic of AuNP-bound (i) and released (ii) DNA labeled with the dyes DL405, FAM, and DL700 of the pH-BOT probe. Excitation wavelengths of 390nm, 480nm, and 670nm, for each dye, respectively. Sodium cyanide dissolution of the AuNP gives rise to the fluorescence recovery of the previously quenched DL405 and FAM, whereas the unquenched DL700 maintains fluorescence. (B) pH dependent fluorescence intensity spectra of DL405, FAM, and DL700 from pH 4 to pH 8, with schematic of DNA-bound dye fluorescence changes in basic (i) versus acidic (ii) conditions. (C) pH dependent ratio of FAM:DL405 with sigmoidal fit (red curve) giving the pH calibration equation. (D) The fluorescence intensity ratio of DL405:DL700 displaying pH independence between pH 5-8. Error bars indicate standard error.
with constant ionic strength (150 mM K\(^+\)) (Figure 3.3B). The sigmoidal pH calibration curve generated from the photoluminescence from pH 4 to 8 is shown in Figure 3.3C. From the graph of fluorescent intensity ratios \textit{versus} pH, the standard curve is plotted in Figure 3.3C and the pH calibration equation is generated (Eqn. 3.1):

\[
I_{\text{FAM/DL405}} = 0.35 + 1.71 \div \left[ 1 + e^{\left(\frac{6.38 - pH_i}{0.41}\right)} \right]
\]

(Equation 3.1)

Where \(I_{\text{FAM/DL405}}\) is the fluorescence intensity ratio of FAM to DL405, and \(pH_i\) is the intracellular pH.

### 3.3.4 Dynamic Monitoring of Intracellular pH

Dynamic monitoring of the cellular uptake and endo-lysosomal processing of the pH-BOT probe was investigated using Lipofectamine2000\textsuperscript{®} assisted transfection of CHO cells following literature procedures.\textsuperscript{132} The CHO cells were grown at 37°C in 10% CO\(_2\) using low glucose media with 10% serum, plated at a seeding density of 30,000 cells per cm\(^2\) in glass bottom 96-well optical plates that were coated with poly-D-lysine for enhanced cellular adhesion. Cells were grown to \~60% confluence and the culture media was exchanged with FluoroBrite\textsuperscript{®} imaging media prior to transfection of the CHO cells with 2.6 nM pH-BOT. The cells were incubated with the transfecting material for 1.5 hours at 37°C in 10% CO\(_2\) prior to media washing, to remove any unincorporated material. This step was carried out to ensure that continuous uptake events did not convolute the observations of the kinetics of the DNA release event and the active reporting of intracellular pH by the pH-BOT.

Cellular internalization of the pH-BOT was verified by collection of high resolution confocal z-stack images of transfected cells, fixed at 2 and 6 h time points with CellMask\textsuperscript{TM} Orange plasma membrane stain. It can be seen in Supporting Figure 3.4 that the pH-BOT fluorescence is localized within the plasma membrane boundary, verifying that the fluorescence response of the pH-BOT can indeed be correlated to the intracellular pH experienced by the probe, and is not an artifact of membrane aggregation.

In Figure 3.5, the dynamic events of endosomal uptake and maturation are monitored by measuring the time dependent changes (1.5 h to 24 h) to the emission spectra for DL405 (blue), FAM (green), and DL700 (red), using live cell confocal fluorescent microscopy. Cellular health and morphology, as well as relative localization of the pH-BOT, are observed by referencing the differential interference contrast (DIC) images. The dye labels are observed to be colocalized in
Figure 3.4 Confocal z-stack images of CellMask Orange® plasma membrane stained CHO cells at 2h (A) and 6h (B) post-transfection with the pH-BOT. Arrows show cellular internalization of the pH-BOT, as the pH-BOT fluorescence is observed most clearly at the focal plane corresponding to center of the cell (ii), as opposed to the outer membrane (i) where the signal is out of focus. Some punctate fluorescent packages are observed to even localize within the base of the cells (iii).

punctate packages indicative of endosomal encapsulation following transfection. The DIC channel allows for the visualization of cell morphology, showing no significant impact on cell health throughout the experiment, as well as apparent uptake of dark, AuNP containing, liposome packages. Inspection of the time dependent changes in the blue, green, and red fluorescence channels show the intracellular processing of the pH-BOT in time.

The intensities of the dyes observed in the selected images (Figure 3.5) and observed though continuous imaging via the tiled overlay images (Figure 3.6 fluorescence and DIC composite images and Figure 3.7 ratio map with DIC overlay images) report on the pH-BOT location and pH evolution, allowing a correlation with the endosomal maturation process. The red channel (DL700) is not quenched by the appended AuNP. A plot of the intensity as a function of time is extracted from images collected every 0.5 h to reveal the DL700 intensity is
Figure 3.5 (A) Fluorescence images of the cellular uptake and processing of pH-BOT probe in CHO cells from 1.5 to 24 hours, following Lipofectamine2000® transfection and media washing at 1.5h. Grey (DIC) images allow for visualization of cell morphology, blue (DL405) reports DNA release from AuNP, green (FAM) indicates intracellular pH after DNA release, red (DL700) shows pH-BOT construct tracking throughout the experiment, and the composite (Overlay) image illustrates colocalization of fluoresce signals within the cells. (B) Pseudo-colored thermal plots of the ratiometric fluorescence of FAM/DL405 indicates intracellular pH (4.5 = purple to 7.5 = red) throughout cellular processing of the pH-BOT. Scale bar: 25 µm.
Figure 3.6 Cell uptake and processing of the pH-BOT probe shown in DIC overlay images of the 3 color confocal fluorescence images with DL405 (blue), FAM (green), and DL700 (red). Scale bar: 10 µm.
Figure 3.7 Cell uptake and processing of the pH-BOT probe shown in DIC overlay images of thermally colored Ratio map of FAM:DL405 fluorescence intensities, with warm colors representing more basic pH environments experienced by the probe and cool colors representing more acidic pH environments (i.e. red to blue = 7.4 to 4.5). Scale bar: 10 µm.
nearly time independent (Figure 3.8A). The observation of nearly continuous DL700 intensity indicates efficient endosomal packaging of the transfected pH-BOT occurs with minimal exocytosis of the packages during the experimental measurements.

![Image](image.png)

**Figure 3.8** Average microscope fluorescence intensity ratios of (A) the change in intracellular DL700 intensity from initial intensity, showing little change over time, and (B) the relationship of intracellular FAM to DL700 intensity, showing decreasing FAM intensity and consistent DL700 intensity over the 24 h time course of the experiment.

The blue (DL405) and green (FAM), relative to red (DL700), fluorescence channels report the cleavage of the pH transponder DNA duplex from the AuNP surface and the evolving pH in the endosomal package as the endosome matures. The release event is signaled by the turn-on of emission for FAM and DL405 caused by loss of the effective SET proximity quenching of the fluorophores’ emission when bound to the AuNP. The turn-on event follows cargo release as the DNA duplex and appended fluorophores diffuses away from the AuNP surface. As observed in the composite image tiles, correlated turn-on of the green and blue emission is observed simultaneously in the cell, which indicates duplex DNA release is occurring within the earliest image frame.

A plot of the intensity of DL405 relative to DL700 (Figure 3.9A) allows the dynamics of the DNA release from the AuNP to be measured. The half-life for release is 1.5 h (red sigmoidal curve in Figure 3.9A) with an observed slow depletion of DL405 intensity (blue line) due to endosomal leakage of the duplex DNA from the endosomal packages.
Figure 3.9  (A) pH-BOT DNA release rate from AuNP core as assessed by the fluorescence intensity ratio of DL405 (SET quenched) to DL700 (unquenched), over the 24 h experiment. Initial release kinetics (▲) plotted to a sigmoidal function curve (red fit) from 0 to 10.5 h. Fluorescence ratio decline (▼) plotted to a linear decay (blue fit) from 11 to 24 h. (B) Calculated average endosomal pH experienced by the AuNP pH-BOT probe during 24 hour cellular processing following Lipofectamine2000® transfection and media washing at 1.5 hours (black line). Shaded boxes represent typical pH values of early endosomes (red: pH 7.4-6), late endosomes (yellow: pH 6-5), and lysosomes (blue: pH 5-4.5). Error bars represent standard deviation. (C) Populations of endosomes at given pH values for selected timepoints over the course of the experiment. Shaded boxes represent typical pH values of early endosomes (EE, red: pH 7.4-6), late endosomes (LE, yellow: pH 6-5), and lysosomes (LY, blue: pH 5-4.5).

After 10 h post transfection, the ratio of DL405 to DL700 begins to slowly decay for the remainder of the experiment with a loss of approximately one fluorescence unit per minute, with a linear decay slope of -0.0177 l/h. Similar results were observed for DNA therapeutic delivery from an AuNP into live cells in an earlier publication. Since the red DL700 channel depicts the location of the pH-BOT throughout the experiment, the observed colocalization of the DL405 and FAM with DL700 implies the lack of DNA damage from enzymatic cleavage of the duplex.
DNA strand during the active visualization of trafficking of the labelled DNA probe sequence in the cell over the course of the experiment.

When evaluating the ratiometric intensities of the pH-BOT relative to its ‘‘always on’’ DL700, it is important to note that within the cell, the intensities of DL405, FAM and DL700 are dependent on the state of the pH-BOT (whether DNA is bound or released), its cellular location (whether endosomally encapsulated or cytosolically diffused), the endosomal environment (EE versus LE, or neutral versus acidic pH), and the microscope’s detector sensitivity for each dye. To understand the ratio of DL405 (\(I_{405}^{\text{observed}}\)) to DL700 (\(I_{700}^{\text{observed}}\)), the fraction of bound (\(a\)) to released (\(b\)) DNA must be considered. In addition, the fraction of endosomally encapsulated vs cytosolically diffuse must also be taken into account. For instance, when \(m\) molecules of labeled DNA escape from the endosome, both the DL405 intensity (\(I_{405}^{\text{observed}}\)) and DL700 (\(I_{700}^{\text{observed}}\)) are reduced from detectability by \(m\) number of fluorophores due to cytosolic dilution as the DNA diffuses out of the endosome and becomes no longer detectable. However, for the fraction of bound DNA molecules (\(a\)), only DL700 is emissive (\(aI_{700}^{\text{bound}}\)), while DL405 (\(I_{405}^{\text{bound}}\)) and FAM (\(I_{\text{FAM}}^{\text{bound}}\)) fluorescence are both quenched by the proximity to the gold nanoparticle and therefore undetectable. Thus, one can predict that as the dye-labelled DNA escapes from the endolysosomal system, the ratio of fluorescence intensity for DL405/DL700 will decrease as described in Eqn. 1

\[
\frac{I_{405}^{\text{observed}}}{I_{700}^{\text{observed}}} = \frac{bI_{405}^{\text{observed}}}{aI_{700}^{\text{bound}} + bI_{700}^{\text{observed}}} > \frac{bI_{405}^{\text{observed}} - mI_{405}^{\text{observed}}}{(aI_{700}^{\text{bound}} + bI_{700}^{\text{observed}}) - mI_{700}^{\text{observed}}} \quad \text{Endosomal Entrapment} \quad \text{Endosomal Escape}
\]

(Equation 1)

Where \(a\) is the number of DNA molecules thiol-bound to the AuNPs in an endosome, \(b\) is the number of DNA released from the particle surface but remaining in the endosome/lysosome. And \(m\) is the number of labeled DNA molecules that have escaped into the cytosol. Based on Eqn. 1, as the DNA is released from the gold nanoparticle surface, it is expected that the escape of each labeled NA molecule from the endosome reduces the intensity of DL405 and DL700 in a linear relationship for one molecular unit of fluorescence intensity at a time, due to the limit of detection of cytosolically dilute fluorophores. If one extends the above relationship to be time-dependent, it becomes apparent that a negative slope is anticipated to arise whenever the rate of
escape exceeds the rate of release, and is linear whenever the rates are constant. Indicating that for the observed data, rates of release dominate for times less than 4 h, but after 12 h the rates of escape dominate (Figure 4A). Between ~4 to 12 h post-transfection the changing rates of release and escape make it difficult to determine if there is a true equilibrium at any particular time. Comparison of the FAM channel to the DL405 indicates that a decrease in the intensity of FAM occurs in time. The decrease is also observed relative to the DL700 (Figure 3.8B). As expected the decrease in FAM intensity relative to DL700 is sharper than the decrease of DL405 relative to DL700 because of the greater pH-sensitivity of FAM. Plotting the decrease in intensity of FAM relative to DL405 can therefore be interpreted in terms of the pH of the endosome using the calibration curve from Figure 3.3C. A quantitative measure of the intracellular pH experienced by the pH-BOT can be assessed by calculation of the endosomal pH from the ratiometric fluorophore intensities extracted from the live-cell fluorescent microscopy images collected every 0.5 h (Figure 3.6). In Figure 3.9B, the average endosomal pH in time is plotted and assigned to values correlated with reported pH ranges for EE, LE and LY environments. From the time-averaged plot it can be seen that the ensemble average endosomal pH experienced by the probe duplex DNA does not acidify to the typical classification of a LY (pH 5.0-4.5) over the course of 24 h, as expected, but rather appears to stall at a pH hovering between early and late endosomes, near pH 6 (Figure 3.9B). An attempt to replicate the potential pH altering effect of the pH-BOT construct, outside of the endosome, was conducted in cuvette using buffers and cell lysate. Room temperature McIlvaine’s buffer in a pH range from 3.9-7.4 showed no significant buffering effect when exposed to the pH-BOT construct over 24 h (Figure 3.10A). To more closely replicate cellular conditions, CHO cell lysate was extracted, and pH adjusted to 5 and 7, prior to exposure to the pH-BOT and incubation the physiological temperature of 37ºC. A slight alkalizing effect was observed for the cell lysate solution at neutral pH, where a solution of pH 7.0 cell lysate gave an average measured pH response of 7.61 ± 0.15, over the course of 24 h at 37ºC. But this effect seemed to be overcome at higher acid concentrations, where a solution of cell lysate acidified to pH 5.0 gave an average measurement of 5.07 ± 0.21 in the presence of the pH-BOT, over the 24 h experimental time frame, (Figure 3.10B). Additionally, the DNA release from the pH-BOT was investigated in cuvette in buffer versus extracted cell lysate, and both conditions demonstrated a much slower rate of release than observed in the live cell experiment (Figure 3.10C&D versus Figure 3.9A) where only about 50% of full release was observed within
Figure 3.10 Cuvette studies of pH-BOT buffering and release in McIlvaine’s buffer and CHO cell lysate. (A) pH-BOT calculated pH from the fluorescence ratio of FAM to DL405 at 0h (blue diamonds) and 24h (red squares) after RT incubation with pH 3.9-7.4 McIlvaine’s buffer solutions. (B) And calculated pH response of the pH-BOT in CHO cell lysate, pH 5 and 7, at 0h (blue open circle), 12h (purple closed triangle), and 24 h (red banded square) incubated at 37°C. (C) Nano-cargo release of the pH-BOT DNA from the AuNP in (C) McIlvaine’s buffer at pH 7.4 and 5.1, and in (D) CHO cell lysate at pH 7 and pH 5, both incubated at 37°C for 24 h.

24 h at 37 °C, demonstrating that buffer and cell lysate do not adequately recreate the endosomal conditions that promote nano-cargo release. Interestingly, Figure 3.10D shows the pH-BOT experienced slower release within the cell lysate solution at low pH, implying that pH is not the driving factor for nano-cargo release inside cells, and may support theories of thiol-containing
cellular proteins and peptides aiding in the displacement of the thiolate-AuNP bond.\textsuperscript{213,214} This further establishes the complexity of the endosomal microenvironment experienced by the pH-BOT and the need for \textit{in vitro} analysis of NP therapeutic fate and processing.

Microscope images of the time dependent pH response, represented as a pseudo-colored thermal map in Figure 3.5B, reveals the expected cellular heterogeneity of the pH-BOT encapsulated in vesicles at various points of maturation within each cell. In Figure 3.5B the red punctate spots indicated an approximate endosomal pH near 7.5, with gradation down to blue spots which represent acidic pH values near 4.5. While the range of colors (pH values) are visible at each time point, early time points are dominated by more neutral (red) endosomes, and later time points show an increased population of lightly acidic (orange-green) endosomes, comparatively. Inspection of Figure 3.5B shows the expected acidification of the endosomes as they mature from early to late (Figure 3.1).

### 3.3.5 pH Population and Packaging Distribution

The average pH experienced by the pH-BOT can be deconvoluted to report the distribution of endo-lysosomal populations in time. In Figure 3.9C, the peak distributions are predominantly observed to proceed from near neutral pH ~6.8 at early time points towards more classical EE and LE pH values at 6 to 5.5, with relatively few endosomes maturing to LY (pH 5-4.5), and the overall distribution maintaining an average pH near 6. The observed population statistics in Figure 3.9C show the endosomal population distributions are not Gaussian. Observation of the non-statistically uniform population profile paired with the asymptotic behavior for the average pH over a 24 h period suggests the possible occurrence of endosomal stalling. Endosomal stalling has been observed for transfection by polymeric and other ionizable transfection agents and been attributed to buffering inside the endosome.\textsuperscript{172,215} Previously, it was suggested that such a behavior may explain the long-term effectiveness of siRNA delivery from an AuNP surface, since siRNA is known to degrade rapidly in endosomes when delivered by a non-protecting transfection agent alone.\textsuperscript{100} The buffering of the endosome does not imply an absence of maturation for the EE to LE and LY, but rather only that the average pH is affected within the endosomes. Alternatively, the observation of secondary populations of higher pH endosomes may represent distributions of packages which have begun to be processed into recycling endosomes for exocytosis. In addition, it is possible that since DNA release continues
to occur up to 3 h post transfection, the reported distribution of pH values may be affected by the presence of unreleased DNA-bound fluorophores within the endosomes, or that the freed DNA cargo is escaping the endosomal pathway prior to LY packaging, convoluting live pH reporting.

**3.3.6 pH-BOT Endo-Lysosomal Colocalization**

In order to evaluate the state of endosomal maturation and determine the intracellular localization of the pH-BOT probe before and after maximal DNA release (2 h and 6 h, respectively), analysis of fluorescence intensity colocalization of the unquenched DL700 from the pH-BOT probe was preformed using commercially available endosomal and lysosomal tracking labels. Endosomal trackers consisted of red fluorescent protein (RFP) conjugates to Rab5 (EE membrane protein) and Rab7 (LE membrane protein), and the lysosomal tracker used was LysoTracker™ DND-99, purchased from Molecular Probes. Figure 3.11 (A-F) displays the cross-correlation functions (CCF) of colocalization for the RFP fluorescence channel versus the DL700 fluorescence channel averaged over five image fields for each time point and tracker, to give a clear picture of the overall colocalization of the pH-BOT with each vesicular component of the endo-lysosomal pathway, before and after complete release of the DNA from the AuNP. The van Steensel CCF is defined by the best Gaussian fit of the plot created by laterally shifting the image pixels in one fluorescence intensity channel “A” (DL700), versus another potentially colocalizing fluorescence intensity channel “B” (RFP) and comparing the effect of the pixel shift on the Pearson’s correlation coefficient ($r_p = \frac{\Sigma (A_i-a) \times (B_i-b)}{\sqrt{\Sigma (A_i-a)^2 \times \Sigma (B_i-b)^2}}$)

where $i$ denotes intensity values for a given pixel, and $a$ & $b$ denote the intensity averages over the whole image per channel, A and B) for each image overlap position.\(^{149,152}\) Here the observation of maxima at the 0 pixel-shift position indicates colocalization of the pH-BOT probe with the EE, LE and LY trackers (Figure 3.11A-F). Slight lateral favoring of maxima away from 0 pixel-shift, for EE and LE trackers indicate a slight degree of partial colocalization, which is most likely due to the slightly larger and more heterogeneous morphologies of EEs and LEs, as compared to LYs.\(^{216}\) This in contrast to an observation of minima at 0 shift, a sign of delocalization, which is not seen for any of the tracker labels compared with the DL700.\(^{149}\) This indicates that the pH-BOT probe is found in all components of the endo-lysosomal pathway at 2 and 6 h after transfection.
Figure 3.11 Endo-lysosomal pathway colocalization analysis depicting (A-F) the cross-correlation functions of DL700, from pH-BOT, with RFP trackers, for early endosomes “EE” Rab5-RFP (A & D), late endosomes “LE” Rab7-RFP (B & E), and for lysosomes “LY” LysoTracker™ DND-99 (C & F), at 2 and 6 hours post transfection, respectively. Average Manders’ colocalization coefficients for the fraction of DL700 colocalizing with RFP for each vesicle type were found to be 0.035, 0.071, and 0.059 for EE, LE and LY at 2 h, respectively; and at 6 h were found to be 0.068, 0.097, and 0.061, respectively. (G) Chromatographic legend of colocalization between FAM and DL405, from pH-BOT, and RFP from vesicle trackers; and (H) colocalization from digitally zoomed single-cell merged confocal fluorescence microscopy images of CHO cells with the pH-BOT probe and RFP vesicle trackers of either (EE), (LE) and (LY), collected at 6 h post transfection. Arrows indicate vesicles with colocalization of RFP trackers with the pH-BOT in EE (i-iii), LE (iv-vi), and LY (vii-ix), with variation in color due to differences in fluorescent intensities of each fluorophore per endosome. The FAM:DL405 emission ratio gave the following pH values: 6.50 (i), 6.50 (ii), and 6.50 (iii), in EE; 6.04 (iv), 6.25 (v), and 6.39 (vi), in LE; and 6.28 (vii), 5.96 (viii), and 6.33 (ix), in LY, for the selected vesicles at 6 h post transfection. White cell outlines for cells of interest were traced from transmitted light images. Scale bar: 10 µm.
Traditional use of $r_p$ alone to determine the degree of colocalization in a fluorescent microscope image is limited by a necessity to have similar intensity values for each fluorescence channel, where an $r_p$ value of 1 would lend to complete one-for-one intensity matching at each pixel, 0 would be the absence of any colocalization, and -1 would mean opposite intensity matching. Thus $r_p$ values between 1 and 0 are difficult to interpret for most non-ideal systems (i.e. microscope images with background noise, differential quantum efficiencies of fluorescent molecules, or non-homogeneous labeling). In the case of commercially available vesicle trackers, it is not possible to ensure 100% detectable labeling of all vesicles in all cells. This can be seen by the more Gaussian shape of the CCF for the LY tracker (Figure 3.11C (2h) and 3.11F (6h)), which is due to the nature of the dye-based vesicle labeling as compared to the EE (Figure 3.11A & D) and LE (Figure 3.11B & E) trackers which are fluorescent protein-fusion labeled. The EE and LE trackers have RFP constructs that require uptake of a viral transfection agent (BacMam 2.0), cell processing and expression of the RFP-Rab fusion proteins, followed by subsequent successful incorporation into the early and late endosomal membranes. Being a non-trivial process, these trackers give less homogeneous labeling and emission profiles per cell, due to the potential for differential expression in cells based on agent uptake and processing factors, as compared to the LysoTracker™, which is an organic dye that readily crosses the plasma membrane of all cells and fluoresces relatively evenly in the acidic organelles of the cells. This difference in labeling combined with the naturally lower quantum efficiency of a far-red or near IR dye, DL700, give rise to the overall low amplitude of the $r_p$ observed in the CCF plots, and could otherwise be interpreted as an absence of colocalization, if not for consideration of the factors discussed, and the CCF analysis.

Comparison of relative amounts of colocalization of the pH-BOT between vesicle types was conducted by calculation of the Manders’ colocalization coefficients ($M1 = \frac{\sum A_i(when B_i > 0)}{\sum A_i}$) for the fraction of DL700 fluorescent intensity colocalizing with the fluorescent intensity of each RFP tracker (Figure 3.9 caption). Although the use of Manders’ colocalization analysis suffers from similar limitations as $r_p$ with sensitivity to labeling efficiency and intensity matching, it can still be used as an additional relative comparison of colocalization as it mathematically dampens noise of intensity variation. From this measure, it can be said that at 2 h post transfection the pH-BOT probe has predominant colocalization with the LE trackers with an M1 value of 0.071 for
LE, followed by LYs with an M1 of 0.059, versus 0.035 for EE. At 6 h, the colocalization with EEs and LYs shifts to M1 values of 0.068 and 0.061, respectively, but the colocalization with LEs remains high with an M1 value of 0.097. The shift towards higher colocalization with EEs at the later time point may indicate a repackaging event of the pH-BOT agent. Comparing the observed colocalization of the DL700 with the RFP trackers and the measured pH values for endosome populations in Figure 3.9C, a disparity is seen between the expected pH values for those vesicles, as the distributions of pH values would be expected to be more acidic, with the majority of the pH-BOT probes localizing into LEs and LYs. To probe for evidence of a potential pH buffering effect, the colocalization of the EE, LE and LY trackers’ RFP fluorescence intensity was compared to the released pH-BOT probe’s FAM and DL405 fluorescence intensities, in order to measure the pH of the colocalized vesicles after full DNA release at 6 h. It can be seen in Figure 3.11H that the oligonucleotide cargo of the AuNP is present in EEs, LEs, and LYs, at 6 h after transfection, as indicated by the colocalization of fluorescent RFP tags on Rab5 and Rab7 proteins for EEs and LEs, respectively, and LysoTracker™ DND-99 for LYs, with the fluorescence of the FAM and DL405 dyes from the pH-BOT. However, the pH measured by the pH-BOT probe dyes are found to be 6.50 in EEs, 6.04, 6.25, 6.39 in LEs, 6.28, 5.96, 6.33 in LYs, for the selected vesicles (Figure 3.11H). This supports the notion that the pH-BOT probe provides its cargo with some protective effect against lysosomal acidification, as measured LE and LY pH values more closely resemble those expected for EEs or “early” LEs.85,208,209

In all, the fate and processing of the pH-BOT appears to be mostly retained in LEs implying a delay in endosomal maturation for the delivered AuNP pH-BOT. The lack of full maturation is reflected in the ensemble average pH being observed to be above pH 6, which is indicative of release and storage only within early endosomes. Upon analysis of the colocalization of the pH-BOT probe with the stains for each vesicle type, it can be seen that the pH-BOT probe is distributed to all vesicular compartments of the endo-lysosomal pathway. The incongruency is likely due to possible pH buffering effects of the reactive thiolate once liberated from the AuNP surface, potentially paired with a protective effect of the AuNP-DNA packing. The appearance of potential buffering has been observed and debated in other nano-transfection systems, in particular PEI delivery vehicles.172,218 The pH-BOT analysis confirms the buffering
effects which may explain the enhanced stability of nucleic acid therapeutic agents delivered by AuNPs.\textsuperscript{100,102}

### 3.4 Conclusion

Nanomaterials-based therapeutics, targeting the systematic manipulation of cellular metabolites, have been at the forefront of biomedical research for the last few decades.\textsuperscript{219} For therapeutics utilizing nucleic acids (DNA, siRNA, shRNA) understanding the pH evolution during endo-lysosomal processing is critical for therapy design. The study shows that the coupling of AuNPs to dye labelled DNA can allow McSET methods to be employed as a real time active reporter of intracellular events. For pH detection the ratiometric measurement of FAM to DL405 intensity demonstrates that the therapeutic cargo is observed in all stages of the endo-lysosomal pathway, the pH appears to be buffered, and the DNA is not degraded over the 24 h experimental timeframe. The observed buffering is believed to be due to protonation of the freed thiolate following cleavage of the Au-DNA thiolate bond. The release event was observed to mitigate the acidification of early and late endosomes and stall the average endosomal pH near 6. The pH buffering of the evolving endosome may provide a protective environment for the DNA cargo, which released with a 1.5 h half-life at pH 6.5. Colocalization analysis of fluorescently labelled vesicles showed no interruption to the maturation of the endo-lysosomal pathway, only buffering of the endosomal pH (Figure 3.11). Since pH buffering is observed, these findings have important ramifications for medicinal uses of AuNP delivery agents, particularly with respect to therapeutic designs that require acidic activation of an attached medicine or acid-based cleavage of an attached cargo to allow therapy release from the nanocarrier. Being aware of the intracellular conditions which promote cargo release and affect therapeutic efficacy will lead to better nanotherapeutic design and the eventual creation of more effective, dynamic nanomedicines to meet the needs of the individual patient.
CHAPTER 4
DIFFERENTIAL UPTAKE OF CELL PENETRATING PEPTIDE LOADED MULTISHELL QUANTUM DOTS IN DRUG RESISTANT CANCER CELLS

4.1 Introduction

Despite improvements in therapeutic approaches for many types of cancer, the development of multidrug resistance (MDR) remains a significant barrier to improving survival time in patients with many types of aggressive cancers. The 5-year survival rate for melanoma (<20%), mesothelioma (<5%), and gliosarcoma (<10%) are all unfavorable, particularly when diagnosed at late stages.\textsuperscript{220-222} Although these aggressive cancers are often resected, in addition to treatment with various chemotherapeutic agents, the remaining cells inevitably develop chemoresistance resulting in the regrowth of highly aggressive, metastatic cancers.\textsuperscript{223-227}

Therapies that increase drug delivery to tumors, while circumventing drug efflux and other mechanisms responsible for chemoresistance, are critically needed to improve patient outcomes. Recent studies have suggested the acquired drug resistance is coupled to both changes in the surface markers and metabolic pathways within the cells.\textsuperscript{228-231} Unfortunately, initial targeting of cancer surface markers with antibodies to overcome drug uptake difficulties, while effective, has not translated into improved treatments, largely due to changes in surface markers with the evolution of acquired drug resistance.\textsuperscript{232,233} While attempts to overcome the metabolic changes using viral vectors to deliver therapeutics are potentially useful in late stage patients, the immune response induced by viral vectors severely limits their usefulness.\textsuperscript{234,235} Non-viral approaches to overcoming drug resistant pathways include the use of cell penetrating peptides (CPP) coupled to nanoparticles to effectively act as a transfection carrier of chemotherapeutics.\textsuperscript{236} CPPs are derived from small sequence fragments of the protein transduction domain of viral sheath proteins, or human carrier proteins. Packaging of CPPs onto the surface of nanoparticles is an important focus area for packaging of therapeutics into non-viral transfection methodologies.\textsuperscript{237-240} When the nanoparticle – loaded with a therapeutic agent – is combined with a CPP, cellular uptake and delivery of the therapeutic agent can be enhanced. Nano-therapeutic delivery agents have also been shown to enhance the survivability and therefore effectiveness of the delivered nucleic acid agent due to reduced nuclease activity, continuous agent release, and pH clamping of the endosomal package.\textsuperscript{56,140,241}
There are reports suggesting that CPPs exhibit selectivity for different cell types. Thus, we have hypothesized that CPPs may also exhibit specificity for chemoresistance in different cancer cells and have designed an approach to screen CPPs for nanocarrier uptake in a cancer cell types with and without chemoresistance.

**Figure 4.1** (A) Schematic diagram of CdTe@CdSe@CdS@ZnS multishell QD showing peptide functionalization with CAAKA and hCT(9-32). (B) Graphic of QD-CPP cellular uptake, showing cellular membrane association (i), internalization (ii), and endosomal entrapment (iii) versus cytosolic distribution (iv).

This study investigates the cellular uptake of various CPP coated 4 nm multi-shelled CdTe@CdSe@CdS@ZnS quantum dots (QDs) (Figure 4.1) and the effective enhancement of cellular internalization in a series cancer cell lines of human melanoma (A375), human mesothelioma (MSTO), and human (LN18, U87) and rat glioma (9L), along with drug resistant
sub-clones. QDs are used for this study as they are ideal optical probes for molecular tracking in cell uptake studies owing to their brightness and resistance to photo-bleaching, and for their comparability to other clinically relevant, non-emissive, nanomaterials. Although the heavy metals that comprise QDs and allow for the unique optical properties, such as cadmium and tellurium, are potentially cytotoxic and therefore not applicable to potential clinical treatments, the use of a non-therapeutic photoluminescent multi-shelled quantum dot (QD), composed of CdTe@CdSe@CdS@ZnS, allows for quantification of nano-package cellular uptake through fluorescence microscopy. The studied CPPs are appended to the QD surface via the N-terminal cysteine which allows for bidentate coordination using the thiol and amine of cysteine to bind onto the ZnS outermost QD shell. A biphasic ligand exchange allows the displacement of the weakly coordinated organic passivating ligands from the as-synthesized QDs, and replacement by the aqueous CPPs. The results of the study easily translate to clinically applicable nanomaterials such as polymer nanoparticles, nanooxides, nanometals, or other nanocarriers.

4.2 Materials and Methods

4.2.1 Multi-shelled QD Synthesis

Spherical 4.3 nm multi-shelled CdTe@CdSe@CdS@ZnS quantum dots (QD) were prepared using a multistep step core-shell synthesis where the Cd stock solution (0.05 M) was prepared by dissolving CdO in tetradecyl phosphonic acid (0.125M) and octadecene (ODE) at 300˚C under N2. The 0.1 M Te stock solution was prepared by dissolving Te in tri-octyl phosphine (TOP) and dilution by an equal volume of ODE. Similarly, 0.1 M solutions of TOP/ODE:Se, TOP/ODE: Cd(acetate), TOP/ODE: S, and TOP/ODE: ZDC (zinc diethyldithiocarbamate) were prepared as multi-shelling precursors. As a special note, the ZDC precursor decomposes at 70°C which acted as both the zinc and sulfur source for the final ZnS shell. For this reason, care must be used during preparation and storage of the precursor, and the powder was dissolved by sonication in an ice bath.

The core CdTe was prepared by microwave heating of the Cd stock (4 mL, 0.05 M) to 220°C at 300 W under N2, followed by rapid injection of the above mentioned Te stock (2 mL, 0.1 M). The solution was maintained at 220°C (300 W) for 2 min resulting in formation of the 3.4 nm CdTe core. Formation of a CdSe interface layer was accomplished in a step-wise SILAR approach, without isolation of the CdTe core by addition of 1 mL TOP/ODE:Se (0.1 M) at 150°C
(300 W) for 2 min, followed by precipitation using toluene and methanol, and centrifugation to isolate the CdTe@CdSe QD. The final two shell layers for the multilayer QD were added by dissolution of the CdTe@CdSe QD into ODE followed by heating to 150°C in the MW cavity (300W), rapid injection of 0.1 M cadmium acetate in TOP/ODE followed by 0.1 M sulfur powder in TOP/ODE. The reaction was cooled to 135°C, and 0.1M zinc diethylthiocarbamate in TOP/ODE was added and allowed to react for 15 min with a final annealing step at 180°C for 15 min. The multi-shelled QD was isolated by precipitation through the addition of MeOH, washed three times by toluene/MeOH, and centrifuged to isolate a free-flowing powder.

The isolated QDs were analyzed by Transmission Electron Microscopy (TEM) using a JEOL-2010 microscope operated at 200 kV. The samples were dispersed on 400 mesh holey carbon grids (Electron Microscopy Sciences) from a toluene solution. The core of the QD consisted of a ~3.4 nm diameter CdTe nanosphere, with half monolayer of CdSe, one monolayer of CdS, and one monolayer of ZnS. The as-prepared QDs were passivated by Tri-octyl phosphine (TOP). The resulting multi-shelled QD had a 5% size dispersity based on TEM analysis. The first exciton for the QD was found to be at 606 nm with the band-edge emission observed at 665 nm (FWHM = 24 nm) with a QY of 33.7%.

4.2.2 Peptide Surface Passivation

Biphasic place exchange reactions were carried out to replace the organic soluble TOP passivating molecules with the hydrophilic peptides through formation of a bidentate coordination to the N-terminal cysteine of the CPPs, as described previously. The purified N-terminus cysteine modified CPP sequences were purchased from RS Synthesis (Louisville, KY). The sequences are available in Table 4.1. The biphasic ligand exchange was accomplished using phase transfer from tetrachloroethylene (TCE) into water, by dissolution of the QD (5 mg, 2 µM) in 2 mL TCE at 60°C. A N2 sparged, aqueous solution (2 mL) containing 2 M peptide and 0.05 M TCEP was added to the TCE solution of QDs. The biphasic reaction was stirred at 60°C for 10 min, followed by dropwise addition of 100 µL of tetraethylammonium hydroxide (20% wt) to deprotonate the terminal carboxylic acid and provide better aqueous solubility in DI H2O, every 10 min until the QDs were fully exchanged, typically within 1 h. Afterwards, the reaction was allowed to phase separate and the water solubilized QD fraction was removed and stored at 4°C. Prior to use, the CPP- QDs were isolated by two successive ethanol precipitations, or by three
successive solvent exchanges of 0.2 µM filtered H2O using a 3k molecular weight cutoff spin filter (VWR), and resuspended in 0.2 µM filtered H2O.

4.2.3 Human Cell Lines and Culture

The human cancer cell lines used in this report were obtained from the American Type Culture Collection (ATCC). The human mesothelioma cell line MSTO-211H (CRL-2081) and the human malignant melanoma cell line A-375 (CRL-1619) A375 skin cancer cells were acquired from ATCC and cultured at 37°C and 5% CO2 in RPMI 1640 medium (Sigma Chemical Co., St. Louis, MO) supplemented with 10% fetal bovine serum (Sigma) and penicillin/streptomycin (Gibco). Drug resistant cells lines were established as described previously. Additionally, human drug resistant glioblastoma lines U-87 MG (ATCC HTB-14) and LN-18 (ATCC CRL-2610) were acquired and cultured at 37°C and 5% CO2 in Dubelco’s Modified Eagles Medium (DMEM 7777, Sigma), with 10% FBS (Sigma), penicillin/streptomycin/amphotericin B (Sigma) and gentamicin (Gibco).

4.2.4 Rat Gliosarcoma Cell Lines and Culture

Rat gliosarcoma cells (9L) were acquired from the neurotissue bank at the University of California, San Francisco. Sub-cultures of 9L cells were expanded in 75 cm2 plastic flasks (T-75 cm2, Nunc) with DMEM (D7777 Sigma Chemical Co., St Louis, MO), supplemented with 2 mM L-glutamine, 100 µg/mL gentamicin, 0.1% pen/strep and 10% fetal bovine serum (HyClone) generating DMEM complete medium. The cultures were incubated at 37°C in a humidified atmosphere containing 5% CO2. The 9L cells were split into 4 identical flasks. Control, BCNU-sensitive 9L cells (9L) were grown as described above. BCNU-resistant 9L cells (9L-R) were created by challenging 9L cells with increasing concentrations of BCNU (10-225 µM) over time. Surviving cells were grown to confluence and then split into new flasks, allowed to reach 80% confluence, and treated again with complete media or complete media containing BCNU.

Responsiveness of 9L and 9L-R sub-cultures to chemotherapeutics was determined using the In Vitro Toxicology Assay Kit (Sigma-Aldrich TOX-6) based on the sulforhodamine blue (SRB) method for monitoring cell viability. Briefly, cells were plated in 96-well plates, in triplicate, at a density of 1x105 cells/mL and treated for 72 h using the range of indicated concentrations of drugs. After 72 h, cells were fixed by adding 50 µL of ice cold 50%
trichloroacetic acid (TCA) per well for 1 h at 4°C. After washing, plates were air dried before addition of 100 µL of 0.4% sulforhodamine blue. The plates were then washed with 1% acetic acid and air dried. The incorporated dye was solubilized by the addition of 200 µL of 10 mM Tris buffer per well. Absorbance was read on a BioRad Benchmark Microplate reader at 490-530 nm to characterize protein concentration in each well.

4.2.5 Cellular Uptake Studies and Fluorescent Microscopy

For each cancer cell line, cells were plated at 30% confluence (~30,000 cells/cm²) on 1.0 coverglass in 24-well plates (Corning), and grown for 24 h at 37°C in 5% CO₂ prior to transfection with 2 pmol of QD-CPP per well (1 nM). After 24 h incubation with the QD-CPPs at 37°C in 5% CO₂ the media was removed and the cells were rinsed with prewarmed PBS, fixed with 4% paraformaldehyde (PFA) (Sigma), stained with DAPI nuclear stain (Life Technologies), and mounted to slides with Fluorogel anti-fade mounting media (Electron Microscopy Sciences), according to the manufacturers’ instructions. Fixed slides were stored at 4°C in the dark when not under microscopic observation.

Cellular uptake of QD-CPPs were analyzed under fluorescent microscopy using a Nikon Eclipse TE2000 inverted microscope. Excitation of QD-CPPs were performed using the Texas Red filter cube (Nikon) with a 500 ms exposure time. The DAPI nuclear stain was used to correct QD fluorescence per cell number, and was excited with a DAPI filter cube (Nikon) for 300 ms of exposure. Differential interference contrast (DIC) images were collected to visualize the cell boundaries, with 35 ms of light exposure. All images were analyzed in ImageJ software for the total QD fluorescence intensity corrected for cell number and quantum yield differences between QD-CPPs, as observed during photoluminescence characterization in cuvette.

4.3 Results and Discussion

4.3.1 Cell penetrating peptides and uptake

The ability of CPP-labelled nanoparticles to utilize endocytotic pathways and bypass multidrug resistance efflux pumps make them an attractive choice for drug delivery. Figure 4.1 shows a schematic of the CPP labelled multi-shelled QD used to assess the cellular uptake dependence for drug resistant cells relative to drug naive cells. The selected protein derived cell penetrating peptides sequences (Table 4.1) were selected from literature and
modified to have a cysteine at the N-terminus for functional attachment to the ZnS surface of the multi-shelled QD, and to allow for biphasic ligand exchange with the as-synthesized organic capping ligands. As shown previously, the cysteine strongly coordinates to the surface through a metal ligand bond.\textsuperscript{101}

The general cellular uptake mechanism of CPPs is highly debated and appears to be sequence and cell-surface specific, with no verified predictability established in the literature.\textsuperscript{92,250,251} The CPPs in Table 4.1 were chosen to explore the uptake behavior and selectivity for drug resistant versus naive cell lines for high charged peptides using Herpes VP22 (+6) and HIV-1 TAT (+9), compared to the low and neutral charges found in the CPP sequence for hCT(9-32) (+1), K-FGF (+1), Ku70 (+0), integrin β3 (+0), and HIV-1 gp41 (+0). The CPPs are co-loaded onto the QD surface at a 9:1 ratio with a short synthetic peptide sequence, CAAKA, which acts as a space filling peptide to better passivate the QD surface while maintaining loose packing of the target CPP sequences, thanks to the short sequence length of CAAKA versus the other CPPs of Table 4.1. The looser packing of co-loading also allows the target CPP sequence to have more visibility to the cell, allowing for better sequence recognition at the cell surface, as well as reducing the overall surface charge that leads to insolubility with the highly charged peptides.\textsuperscript{252}

Three of the selected peptides in Table 4.1, HSV1-VP22, HIV-TAT, and HIV-gp41 (Table 4.1), are derived from segments of viral proteins that have an added cysteine mutation at the N-terminus. VP22 is a truncation of the tegument protein, viral protein 22 (VP22) found in herpes simplex virus (HSV-1).\textsuperscript{253–255} HIV-TAT and HIV-gp41 are derived from subunits of viral capsid proteins of the human immunodeficiency virus type-1 (HIV-1). HIV-TAT is a fragment of the transactivator of transcription (TAT) protein and HIV-gp41 is derived from the HIV glycoprotein 41 (gp-41).\textsuperscript{30, 38} These three viral peptides have successfully been utilized to transport cargo across the cell membrane in various cell lines.\textsuperscript{90,254,258–261} Although the exact mechanism of internalization for these viral CPPs is still contested, it is thought that the peptides enter through one of the mechanisms of endocytosis.\textsuperscript{262} The remaining CPPs are based on cysteine modified peptide fragments from proteins identified from the literature that induce cellular uptake. Ku-70 is derived from the Bax-binding domain of the protein Ku-70, which acts in DNA repair inside the nucleus.\textsuperscript{263} The excised Ku-70 peptide sequence is also known as a Bax-inhibiting peptide (Bip), and has demonstrated low toxicity in cells, but the internalization
mechanism is also undetermined.\textsuperscript{264} hCT(9-32) is derived from the C-terminal fragment of the human calcitonin protein hormone, and is also thought to internalize through a mechanism of endocytosis; however, it is suggested that high concentrations are needed to facilitate membrane disruption.\textsuperscript{265–267}

Table 4.1 Table of selected Cell Penetrating Peptides, displaying numerical designation, CPP name, sequence and length, charge and isoelectric point (Pi) at pH 7.4, molecular weight in Daltons, and percentage of hydrophobic residues (L\%). CPP origin is denoted by \( \downarrow \) for synthetic, \( \dagger \) for viral proteins, and \( \ddagger \) for non-viral proteins. (*cysteine mutation, not present in the original sequence.)

<table>
<thead>
<tr>
<th>CPP</th>
<th>Sequence &amp; Origin</th>
<th>Length</th>
<th>Charge, pH 7.4</th>
<th>Pi</th>
<th>MW, Da</th>
<th>L %</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAAKA</td>
<td>CAAKA ( \downarrow )</td>
<td>5</td>
<td>+1</td>
<td>8.97</td>
<td>462.6</td>
<td>75</td>
</tr>
<tr>
<td>HSV1-VP22</td>
<td>C*DAATATGRSAASRPTERPRAPA ( \dagger )</td>
<td>35</td>
<td>+6</td>
<td>12.3</td>
<td>3748.2</td>
<td>40</td>
</tr>
<tr>
<td>HIV-TAT</td>
<td>C*YGRKKRRQRRR ( \dagger )</td>
<td>12</td>
<td>+8</td>
<td>12.4</td>
<td>1663.0</td>
<td>0</td>
</tr>
<tr>
<td>HIV-gp41</td>
<td>C*GALFLGWLGAAGSTMGA ( \dagger )</td>
<td>18</td>
<td>+0</td>
<td>5.33</td>
<td>1683.0</td>
<td>56</td>
</tr>
<tr>
<td>Ku-70</td>
<td>C*PMLKE ( \ddagger )</td>
<td>6</td>
<td>+0</td>
<td>6.28</td>
<td>719.9</td>
<td>60</td>
</tr>
<tr>
<td>hCT(9-32)</td>
<td>C*LGTYTQDFNKFHTFPQTAIGVGA ( \ddagger )</td>
<td>25</td>
<td>+1</td>
<td>7.1</td>
<td>2714.1</td>
<td>40</td>
</tr>
<tr>
<td>integrin ( \beta )3</td>
<td>C*VTVLALGALAGVGVG ( \ddagger )</td>
<td>16</td>
<td>+0</td>
<td>5.33</td>
<td>1399.7</td>
<td>63</td>
</tr>
<tr>
<td>K-FGF</td>
<td>C*AAVALLPAVLLAHLLAP ( \ddagger )</td>
<td>18</td>
<td>+1</td>
<td>7.15</td>
<td>4533.1</td>
<td>94</td>
</tr>
</tbody>
</table>

The CPPs integrin \( \beta \)3 and K-FGF both derived from fragments of signal sequences of proteins. The integrin \( \beta \)3 peptide sequence is derived from the hydrophobic region of the signal sequence of human integrin \( \beta \)3 protein.\textsuperscript{268} Similarly, the K-FGF peptide sequence originates from the signal sequences of fibroblast growth factor found in Kaposi’s sarcoma cells.\textsuperscript{269,270} Integrin \( \beta \)3 and K-FGF peptide sequences have been observed to enter cells by means of endocytosis, but it is believed that direct translocation across the membrane may also be possible for such highly hydrophobic molecules.\textsuperscript{271–274}
4.3.2 Synthesis & Biofunctionalization of Multishelled Quantum Dots

The multishelled QD (CdTe@CdSe@CdS@ZnS) structure was chosen to provide the targeted color of the QD (λem = 606 nm (full width half maximum 24 nm) and increased stability of the QD in biological media due to the hetero-layering. Figure 4.1 shows a schematic of the multishelled QD used in this study along with the two example peptides attached to the surface through the N-terminal cysteines.

Figure 4.2 Characterization of the multishell QDs with (A) TEM of CdTeSe cores (i), CdS shelled (ii), and ZnS capped (iii) QDs. (B) Size distributions for CdTeSe cores (i), CdS shelled (ii), and ZnS capped (iii) QDs. And (C) absorption and emission spectra for CdTeSe cores (i), CdS shelled (ii), and ZnS capped (iii) QDs.
Briefly, the core CdTe@CdSe particles show absorption and emission features at 562nm and 579nm, respectively. As a shell of CdS is added to the particles, there is a red shift to 568nm and 584nm. Finally, the CdTe@CdSe@CdS@ZnS particles showed an absorption and emission of 601nm and 606nm, respectively. TEM images are shown at each stage of shell addition, along with size distribution as evidenced in Figure 4.2. The UV-VIS shows a strong first excitonic peak 601 nm with an emission peak at 606 nm with a full width half max of 24 nm, displaying a sharp emission profile (Figure 4.2). In addition, the size of the particle (4.2 nm) was chosen to be resolvable from autofluorescence and to allow crossover into downstream studies, as particles under 5.5 nm are able to be cleared in vivo by the renal system. In addition, the thickness of the ZnS shell aides in shielding the cell from potential toxic leeching of the CdTe core as well as providing a stable outer layer for peptide assembly.

Table 4.2 Optical properties of CPP functionalized QDs, showing CPP-dependent Quantum Yield (Φ), First Exciton Absorption Wavelength (λAbs), and Maximum Photoluminescence Wavelength (λPL).

<table>
<thead>
<tr>
<th>QD-CPP</th>
<th>Φ (%)</th>
<th>λAbs (nm)</th>
<th>λPL (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAAKA</td>
<td>11.3</td>
<td>644</td>
<td>670</td>
</tr>
<tr>
<td>HSV1-VP22</td>
<td>7.4</td>
<td>644</td>
<td>668</td>
</tr>
<tr>
<td>HIV-TAT</td>
<td>37.0</td>
<td>632</td>
<td>658</td>
</tr>
<tr>
<td>HIV-gp41</td>
<td>8.3</td>
<td>643</td>
<td>664</td>
</tr>
<tr>
<td>Ku-70</td>
<td>2.3</td>
<td>644</td>
<td>674</td>
</tr>
<tr>
<td>hCT(9-32)</td>
<td>2.4</td>
<td>648</td>
<td>670</td>
</tr>
<tr>
<td>integrin β3</td>
<td>18.4</td>
<td>643</td>
<td>658</td>
</tr>
<tr>
<td>K-FGF</td>
<td>12.4</td>
<td>639</td>
<td>664</td>
</tr>
</tbody>
</table>

Aqueous functionalization of the phosphine passivated QDs, which were organically soluble as-synthesized, required biphasic ligand exchange (Figure 4.3). The selected CPPs (Table 4.1) were coupled to the QD at a loading ratio of 9:1 CPP-X to CAAKA (CPP1), used as a synthetic spacer peptide, to allow for the longer CPPs to avoid tightly packing on the QD surface, as this may hinder sequence recognition by the target cells. A molar excess of 1000:1 peptides to QDs was used to drive the aqueous transfer of the previously organically soluble QDs.
Figure 4.3 Passivation of CdTe@CdSe@CdS@ZnS multishell QDs biofunctionalized with cell penetrating peptides (CAAKA, HSV1-VP22, HIV-TAT, HIV-gp41, Ku-70, Hct(9-32), Integrin β3, and K-FGF). Schematic (A) shows exchange of organic ligands, TOP and TDPA (i), for 10% CPP-X and 90% CAAKA (ii). Image (B) displays QD solution before and after phase exchange in room light (i & iv) and under fluorescence excitation (ii & iii). The absorbance (C) and emission (D) spectra for each QD-peptide sample demonstrate slight changes in the surface environment per peptide.
In a typical biphasic exchange the QD, dissolved in TCE, was observed to transfer to the upper aqueous, peptide-containing layer within 1 h. By increasing the alkalinity of the aqueous phase (pH > 8.0, peptide NH2 pKa ~ 8), the peptide is deprotonated and transports the QD from the organic to the aqueous layer. Neutralization of the aqueous layer and back extraction yielded water-soluble QDs, capable of use in the biological application. The selected screen of CPPs, selected from literature sources (Table 4.1), were intentionally mutated to contain N-terminal cysteine residues to provide an amine and thiol bidentate binding motif to passivate the ZnS outer shell of the QDs. Since luminescence of the QD is directly related to the electronic environment of the semiconducting elements of the material, changes to the surface passivation chemistry led to observable changes in the optical properties of the QDs. Thus, it was seen that passivation with the selected screen of CPPs shifted the absorbance and emission peaks closer to 640 and 660 nm, respectively, with slight variations due to sequence differences of the CPPs (Figure 4.3C & D). As shown in Figure 4.3, the exchange of the peptide onto the QD surface is evidenced by aqueous solubility and a shift in the absorption and emission properties. The absorbance and emission spectral profiles of the QD-CPPs are observed to shift slightly with the changing molecular environment at the surface of the QD due to the unique chemical properties of the loaded CPPs (Figure 4.3C & D). Relatedly, the quantum yields (Φ) for the aqueous QDs is dependent on the identity of the peptide coating as well. The experimental data is tabulated in Table 4.2. The spectral shift and quantum yield is accounted for in the cell studies to allow for more accurate quantification of cellular uptake.

### 4.3.3 Cellular Uptake Studies

The QD-CPP constructs were used to challenge the drug resistant and naive cancer cell lines for a 24 h incubation period and were then washed with PBS, fixed and co-labelled with DAPI nuclear stain, as described above. Fixed cells were imaged using a widefield inverted fluorescence microscope for each QD-CPP treatment (Figure 4.4). Figure 4.4 displays fluorescence images of the blue DAPI nuclear stain and red intracellular QD emission. As well as overlays with the DIC channel to visualize cellular morphology. All cell lines appear healthy at the time of fixation, indicating no toxic effect from the 24 h QD incubation. Red photoluminescence of the QDs within each cell line is visible at varying intensities, depending
Figure 4.4 Fluorescence images of QD-CPP uptake in rat gliosarcoma (9L) and drug resistant (9L-R), human patient drug resistant gliosarcoma (LN18 & U87), human melanoma (A375) and drug resistant (A375-R), and human mesothelioma (MSTO) and drug resistant (MSTO-R), fixed at 24 h with DAPI nuclear stain. Scale bar:100 µm.
on the peptide functionalization and cell line, which is indicative of differential cellular uptake and processing of the QDs being correlated with the identity of the covalently attached CPP.

In addition to uptake differences being peptide specific, individual cell types presented unique QD-CPP fluorescence profiles. This may potentially indicate an effect from differential phenotypic expression of cell membrane surface components between various tissue types and MDR status on the interactions of the CPPs, leading to increased or decreased cellular internalization of the QD-CPP agents. Additionally, the morphological patterns of intracellular QD photoluminescence were observed for each QD-CPP per cell line (Figure 4.4). Small punctate patterns are indicative of endosomal encapsulation of the QD-CPP construct, whereas the observation of diffuse, evenly distributed intensity throughout the cell represents cytosolic localization of the QD-CPP agents.

To quantify QD-CPP cellular uptake and localization in endosomes versus cytosol, the intensity levels of red photoluminescence within the cells were analyzed using Nikon Elements and ImageJ Software. Images were background subtracted and normalized to maximum intensity per cell number, and the extracted fluorescence values were corrected for relative differences in quantum efficiency for each QD-CPP construct, to allow for accurate comparison of cellular uptake. In Figure 4.5, the corrected cellular uptake of each QD-CPP construct is displayed in comparison to naive and resistant cell lines, for melanoma skin cancer (Figure 4.5A), mesothelioma lung cancer (Figure 4.5B), and rat gliosarcoma brain cancer (Figure 4.5C), as well as for both human patient excised MDR gliosarcoma cell lines (Figure 4.5D). Uptake profiles plotted from analysis of the QD intensity in the microscopy data reveals cellular uptake selectivity for the CPPs that are cell line specific. HSV1-VP22, Ku-70, and hCT(9-32) exhibit preferential cellular uptake in the drug resistant A375-R line versus the naive A375 line, with a statistical significance of $p \leq 0.05$ for HSV1-VP22 and hCT(9-32), and $p \leq 0.01$ for Ku-70 (Figure 4.5A). A similar trend was observed for hCT(9-32) and HIV-gp41 in the 9L rat gliosarcoma drug resistant line versus the drug naive cells, with a statistical significance of $p \leq 0.0001$ and $p \leq 0.01$, respectively (Figure 4.5C). And although no direct comparison with drug naive human gliosarcoma cells was able to be obtained for this study, both human MDR gliosarcoma cell lines displayed the highest uptake for Ku-70 and hCT(9-32) (Figure 4.5D). Interestingly, contrasting results were observed for the mesothelioma cell lines where the naive MSTO cell lines showed the opposite trend in uptake compared to the drug resistant line, and
experienced higher QD-CPP uptake of Ku-70 and hCT(9-32), versus the drug resistant line (Figure 4.5B), highlighting the complexity of the CPP-cell surface interaction.

**Figure 4.5** Uptake profiles of the studied QD-CPP constructs in (A) human melanoma (A375) and drug resistant (A375-R), (B) human mesothelioma (MSTO) and drug resistant (MSTO-R), (C) rat gliosarcoma (9L) and drug resistant (9L-R), and (D) human patient drug resistant glioblastoma (LN18 & U87), fixed at 24 h, measured by microscopic fluorescence intensity, corrected for the quantum yield of each peptide-QD conjugate, and normalized for cell number by DAPI nuclear stain counting. Displaying statistical differences of $p \leq 0.5$, 0.1, 0.001, and 0.0001 as *, **, ***, and ****, respectively, for uptake in comparison to 100% CAAKA loaded QDs (black), and drug resistant versus drug naive cell lines (red, in parentheses).
Further studies are underway to understand the nature of the cell-CPP interactions leading to the observed results. Since the QDs were loaded with CPPs at a ratio of 9:1 with the CAAKA short synthetic peptide spacer, occupying 90% of the QD surface sites, to allow for better visibility of the literature-sourced sequences in Table 4.1 to be accessible by the cell surface, analysis of QD-CPP cellular uptake versus the 100% CAAKA loaded QD was performed as a control. It was found that for all cell lines, QDs with 10% loading with Ku-70 and hCT(9-32) showed improved uptake versus CAAKA alone, giving the statistical significance of $p \leq 0.0001$ for all lines except A375 (hCT(9-32) $p \leq 0.001$), MSTO (Ku-70 $p \leq 0.001$ and hCT(9-32) $p \leq 0.01$), and MSTO-R, the latter of which did not show statistical significance above the 100% CAAKA-QD sample for any of the tested constructs (Figure 4.5). Additionally, the Herpes virus derived peptide, HSV1-VP22, showed significant uptake over the 100% CAAKA loaded QDs in 9L & 9L-R ($p \leq 0.001$), U87 ($p \leq 0.01$), and A375-R ($p \leq 0.5$) (Figure 4.5). These data emphasize the complex and somewhat ambiguous nature of CPP cellular targeting and stress the need for a more comprehensive understanding of design considerations for cell-specific CPP targeting approaches to be fully realized.

Much of the recent literature suggests that cellular uptake of CPPs is highly dependent on electrostatic interactions of cell surface markers in addition to the positive charge of the peptide.\textsuperscript{262} Although the set of peptides tested was not exhaustive, this data indicates that uptake of the CPPs could rely more heavily on cellular surface markers than previously anticipated. Drug resistant cancer cell lines are known to express different surface markers depending on the amount of resistance to the chemotherapeutics, however, as most research indicates that CPPs gain access to the intracellular lumen through various endocytic pathways,\textsuperscript{92} these results suggest that CPPs are not only sensitive to cell type, but potentially to changes in the cellular membrane, the surface receptors as well.

### 4.3.4 Cellular Localization Analysis

The CPP modified QDs were analyzed for intracellular particle localization, to probe the question of whether an endosomal versus cytosolic fate for the QD-CPPs is a cell- or CPP-dependent phenomenon. In Figure 4.6A, high resolution confocal z-stack images display cellular localization of internalized QD-CPPs in three-dimensional space, with an example of more endosomal punctate
Figure 4.6 (A) Confocal fluorescent z-stack images of intracellular localization for QD-CPP constructs displaying more punctate endosomal fluorescence for QD-CAAKA (i) and more diffuse cytosolic localization for QD-hCT(9-32) (ii) in 9L cells. (B) Quantification of QD-CPP endosomal versus cytosolic localization by area of punctate fluorescence, showing mostly mixed endosomal (10 µm² single endosome) and cytosolic (900 µm² whole cell) localization for naïve and drug resistant human melanoma (A375 & A375-R), human mesothelioma (MSTO & MSTO-R), rat gliosarcoma (9L & 9L-R) and human glioblastoma (LN18 & U87), with (ii-v) endosomal area range for CAAKA versus the best performers: HSV1-VP22, Ku-70, and hCT(9-32), respectively.
fluorescence shown in for the 100% CAAKA loaded QD control (Figure 4.6A-i), and more
cytosolic diffuse localization for 10% hCT(9-32) loaded QDs (Figure 4.6A-ii), at 24 h post
transfection with the QD-CPP constructs. From these and previous cell images, it can be seen
that all of the QD-CPP constructs display varying proportions of both punctate and diffuse
fluorescence indicating different packing and escape profiles for each cell line that do not seem
to be entirely CPP-dependent (Figures 4.6A and 4.4).

In order to assess cytosolic versus endosomal localization of the QD-CPP packages
within cells, two-dimensional cell images were analyzed for the average fluorescence area, with
small punctate fluorescence areas indicating endosomal localization and large diffuse
fluorescence areas indicating cytosolic QD-CPP distribution (Figure 4.6B). Comparing the
results of Figure 4.6B to the average 2D area of a whole cell (~900 µm²) versus that of a single
endosome (~10 µm²), it is observed that most of the internalized QD-CPPs were at least partially
confined to endosomes with some QD-CPPs having been cytosolically delivered or having begun
to diffuse into the cytosol by the 24 h time point, as the average package size for all CPPs and
cell types is well below the “whole cell” or fully cytosolic value of 900 µm². Taking a closer
look at the average fluorescence package sizes of QD-CAAKA versus the best uptake
performers, HSV1-VP22, Ku-70 and hCT(9-32), it can be seen that the localization profiles seem
to be very cell type dependent (Figure 4.6Bii-iv). The A375 and A375-R skin cancer cells show
greater than 10 µm² average package sizes for all peptides, although package sizes of 20-30 µm²
could be indicative of small endosomal agglomerates. The MSTO and MSTO-R lung cancer
cells show greater cytosolic localization profiles for CAAKA (near 80 µm² or 84±42 and 72±18
µm², respectively) as well as the best performers (~60-130 µm²). This in comparison to glioma
cell lines which more distinctively show endosomal entrapment for the QD-CAAKA samples
(near or below 10 µm²), with HSV1-VP22, Ku-70 and hCT(9-32) each showing more cytosolic
(average size greater than 30 µm²) for one or more glioma cell lines (Figure 4.6Bii-iv). The
experimental observation is an interesting finding as it has previously been assumed that
nanoparticle delivery constructs disassemble during packaging in the endo-lysosomal pathway
allowing escape of the cargo with only the NP being trafficked out of the cell via exocytosis. Determination of the amount of cytosolic delivery versus endosomal escape in the cell, and
what cellular uptake mechanisms may be involved with each of the cell penetrating peptides will
require future studies.
4.3.5 Computed CPP Orientation and Packing Analysis

Much of the recent literature suggests that cellular uptake of CPPs is highly dependent on electrostatic interactions of cell surface markers in addition to the positive charge of the peptide.\textsuperscript{262} Although the set of peptides tested was not exhaustive, the data in Figures 4.4–4.6 suggests that uptake of the CPPs could rely more heavily on CPP interactions with cellular surface markers than previously anticipated. Most research indicates that CPPs gain access to the intracellular lumen through various endocytic pathways,\textsuperscript{92} many of which require receptor mediated binding as an initiation step.

\textbf{Figure 4.7} PEP-FOLD generated secondary structures of the studied CPPs as possible folding orientations of peptides bound to the QD surface \textit{via} the N-terminal cysteine thiol.

Since acquisition of drug resistant status in cancer cell lines is known to cause a change in membrane composition and expression of different surface markers\textsuperscript{281} – depending on the
amount of resistance to the chemotherapeutics – these results may suggest that ability and efficiency of CPP cellular uptake is not only sensitive to membrane surface markers available per cell type, but also potentially to changes in the cellular membrane structure, and prevalence or identity of surface receptors within tissue classes as well. To further understand the peptide dependent uptake behavior, the nature of the potential packing and orientations of the appended peptides on the surface of the QD was analyzed using the PEP-FOLD structure predicting software service,\(^{282}\) to assess the most energetically favorable structural conformations for the CPP sequences under physiological conditions. The generated results were further filtered to select the most plausible QD-bound secondary structure, which allowed for unhindered access of the thiol from the N-terminal cysteine to bind to the QD surface.

**Table 4.3** Calculated peptide properties and loading level on the 4 nm diameter QD (surface area ~12.5 nm²), based on the PEP-FOLD generated CPP structures, displaying folded peptide height and diameter given in Angstroms (Å), binding footprint in square nanometers (nm²), and the maximum number of peptides a 4 nm QD could accommodate for single peptide loading (100%), and ratioed peptide loading (90% & 10%) for CAAKA:CPPX.

<table>
<thead>
<tr>
<th>CPP</th>
<th>Height (Å)</th>
<th>Diameter (Å)</th>
<th>Footprint (nm²)</th>
<th>100% Loading (# per QD)</th>
<th>90% CAAKA (# per QD)</th>
<th>10% CPPX (# per QD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CAAKA</td>
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<td>0.6</td>
<td>79.0</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>HSV1-VP22</td>
<td>31.5</td>
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<td>5.6</td>
<td>9.0</td>
<td>39.9</td>
<td>4.4</td>
</tr>
<tr>
<td>HIV-TAT</td>
<td>27.6</td>
<td>25.6</td>
<td>5.1</td>
<td>9.8</td>
<td>41.6</td>
<td>4.6</td>
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<tr>
<td>HIV-gp41</td>
<td>31.7</td>
<td>15.2</td>
<td>1.8</td>
<td>27.7</td>
<td>60.0</td>
<td>6.7</td>
</tr>
<tr>
<td>Ku-70</td>
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<td>13.1</td>
<td>1.3</td>
<td>37.3</td>
<td>64.0</td>
<td>7.1</td>
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<tr>
<td>hCT(9-32)</td>
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<td>2.2</td>
<td>23.2</td>
<td>57.3</td>
<td>6.4</td>
</tr>
<tr>
<td>Integrin β3</td>
<td>23.1</td>
<td>9.8</td>
<td>0.8</td>
<td>66.6</td>
<td>69.8</td>
<td>7.8</td>
</tr>
<tr>
<td>K-FGF</td>
<td>25.6</td>
<td>19.2</td>
<td>2.9</td>
<td>17.4</td>
<td>52.5</td>
<td>5.8</td>
</tr>
</tbody>
</table>

In Figure 4.7 it can be seen that each peptide adopts a different degree of ordered secondary structure. These differences lead to unique structural morphologies which do not trend with the properties of the primary sequences, for instance longer sequence length peptides do not always display a proportionally greater height from the QD surface than shorter sequence peptides (Table 4.3). Also, the folding orientations alter each peptide’s effective footprint, or the amount of “floorspace” that the peptide occupies on the QD surface, impacting CPP packing efficiency and loading level on the QDs (Table 4.3). Both the factors of CPP orientation and
packing play an important role in the ability of the transduction sequences of the peptides to access and interact with the cellular membrane to promote cellular entry. The maximum possible numbers of peptides able to spatially fit on the QD surface (ignoring more complex steric or repulsive molecular interactions) can be seen in Table 4.3 for 100% loading and 10% loading versus CAAKA. Notably, the co-loading of the short CAAKA filler sequence not only has the ability to aid in visibility of the target CPPs, giving a calculated ~7-16 Å of head clearance (depending on CPP), but may also aid in QD stability by increasing the surface passivation and allowing for more binding sites on the QD to be occupied than if loaded with the target sequences alone (Table 4.3).

4.4 Conclusion

The selectivity of the various CPPs for drug-resistant cancer cell lines was analyzed and compared to the drug-naive cancer lines. The results of the study indicate that the uptake behavior is highly dependent on the cell type and its level of acquired resistance, as well as the CPP. The studied CPPs were selected from reported viral or signal protein sequences that have been demonstrated to cross cell membranes. The CPP sequences studies were modified with a N-terminal cysteine appended to truncated fragments of highly charged viral capsid proteins (HIV-TAT and HSV1-VP22)\textsuperscript{255,257} and protein transduction domains with low charge (hCT(9-32), Ku-70, and K-FGF),\textsuperscript{264,266,283} and neutral transduction sequences (HIV-gp41 and integrin-β3).\textsuperscript{259,268} The CPPs co-loaded onto the multi-shelled QDs at a 1:9 molar ratio versus the short synthetic peptide CAAKA. As indicated in the PEP-FOLD analysis (Figure 4.7) co-loading increases the target CPP availability to the cell surface and potentially enhance recognition of the CPP sequence with the cell surface, by mitigating close packing of the CPP backbones around the QD delivery vector. The results indicate each CPP exhibits cell-specific uptake profiles with unique responses being observed for each cell line and peptide (Figure 4.4 & 4.5). Despite these cell-specific variations, significant uptake is observed for the hCT(9-32), Ku-70 and HSV1-VP22 peptides for the majority of the selected screen of cell lines, with significant increases being observed for the drug resistant human melanoma and rat gliosarcoma cells (p < 0.05 and 0.0001, respectively), which may be due to changes in membrane protein expression as cancer cells acquire drug resistance. Inspection of the sequences reveals the uptake profiles do not appear to be dependent on the peptide charge, length, or secondary structure; although peptide
packing on the QD surface may affect cellular uptake. This work highlights the utility of multi-shelled QDs for effective CPP-mediated uptake screening in multiple cell lines using fluorescent microscopy, as well as demonstrates the non-trivial considerations involved in designing targeted nanotherapeutics for drug resistant cancer treatment.

This work demonstrates that for the selected cells lines, differences in QD-CPP cellular uptake is dependent on the drug resistance status as well as the specific CPP used. Results from this study provide insight into the use of CPPs in cancer lines, as the CPPs can have variable transfection efficiency depending on cell type and drug resistance status of the cells. This is important for future use of drug delivery designs, as targeting a cancer in vivo does not have a “one-size-fits-all” solution, and different CPPs might be needed depending on the drug resistant status of the cells at the time of delivery. It is also observed that the QD does not cause toxicity and provides valuable insights into cell-specific uptake profiles. Future studies are needed to assess the effects of co-appending a drug to the surface of the QD to determine the outcomes of a multi-biofunctional multi-shelled QD-CPP, within these cell lines, in order to better understand the utility of CPP conjugated nanoparticles as a viable therapeutic for drug delivery.
CHAPTER 5
OPTICAL AND MAGNETIC MEASUREMENT OF NANOTHERAPEUTIC DELIVERY INTO CANCER CELLS USING PEPTIDE CONJUGATED MAGNETIC GOLD (Fe3O4@Au) NANOPARTICLES

5.1 Introduction

Utilization of magnetic materials at the nano-scale has found diverse applications in medicine and technology, with superparamagnetic iron oxide (SPIO) being the most abundant and cost effective. Complimentarily, nanoscale gold is relatively nontoxic, easily biofunctionalized, and has a strong surface plasmon resonance able to interact with fluorophore for sensing applications. Layering the advantageous attributes of a magnetic nanoparticle with the biostable and functional surface of a gold nanoparticle, produces an effective "magnetic-gold particle" (MGP) (Figure 5.1).

Figure 5.1 Schematic diagram of a multi-functional “Magnetic Gold” Fe3O4@Au Nanoparticle (MGP), displaying optical and magnetic utility from the Au shell and Fe3O4 core, respectively.
Combining the utility of magnetic and gold nanomaterials, these superparamagnetic gold-shelled iron oxide nanoparticles have numerous biological applications in magnetic and x-ray contrast agents, diagnostics, hyperthermia, and drug delivery. Thus, the scope of therapeutic applications for MGPs is near limitless, with function being determined by design of their bio-conjugation cocktail (small molecule drugs/ligands, proteins/peptides/antibodies, and/or nucleic acids). Additionally, many stabilizing agents bind securely to the surface of gold, and when the Fe3O4 is shelled by the Au top layer, the gold is able to protect not only the biological environment from Fe leeching, but also help mitigate magnetic aggregation, which is a common problem in magnetic nanomaterials.

Superparamagnetic gold-shelled iron oxide nanoparticles have numerous biological applications in magnetic and x-ray contrast agents, diagnostics, hyperthermia, and drug delivery. Thus, the scope of therapeutic applications for MGPs is near limitless, with function being determined by design of their bio-conjugation cocktail (small molecule drugs/ligands, proteins/peptides/antibodies, and/or nucleic acids). Additionally, therapeutic strategies requiring intracellular delivery (such as pharmaceutical or genetic therapy approaches) predicate a need for quantification of nano-therapeutic cellular uptake to determine the efficacy of the treatment. Future clinical applications will also necessitate the ability to track nano-therapy biodistributions in live patients. Utilizing the dual magnetic- and optically-responsive MGP platform allows for analysis of uptake through magnetic measurement as well as via microscopy. This two-fold sensing tool finds great purpose in the diagnosis and treatment of cancers, as some cancers are inherently more accessible for biopsy and visible observation (such as more superficial melanoma), versus others (such as gliosarcoma) which can present deep within delicate tissue, making excision and direct observation a near-impossibility. However, the use of a multifunctional MGP could allow for multiple modes of visualization and treatment, benefitting countless patients.

In order to be a successful diagnostic and treatment platform, the MGP must be able to be efficiently and specifically taken up by the target cells and tissues. Facilitation of efficient cellular internalization can be accomplished through appendage of cell penetrating peptides (CPPs) which have been isolated from a variety of sources, including protein transduction domains of viral (HIV-TAT: the transactivator of transcription for the human immunodeficient virus) and signaling (Ku-70: a Bax inhibiting peptide) proteins, as well as peptide hormones.
An added advantage of CPP co-assembly is their observed cell-selective internalization, which could allow for cell-type specific targeting. Direct delivery of therapeutics to tumors is of great need since most existing anticancer agents demonstrate systemic distribution that limits their therapeutic efficacy and generate undesirable side effects. The implementation of MGP-CPPs as efficient and specific therapeutic carriers in medicine could reduce the systemic dispersal of cytotoxic anticancer drugs, and reduce the dosage required through more localized targeting – all while providing a multi-traceable delivery agent to track therapeutic delivery and bioaccumulation – overall improving patient health and quality of life.

In this study MGPs were synthesized and bio-functionalized with a small molecule as drug mimics, as well as a small screen of cell penetrating peptides (CAAKA, HIV-TAT, Ku70, and hCT), and observed in human melanoma (A375) and rat gliosarcoma (9L) cancer cell lines via optical microscopy, superconducting quantum interferometry, and magnetic resonant imaging in a tissue mimic. Moreover, with the help of the CPPs cellular targeting and internalization was accomplished. Cellular uptake was quantified and compared across the magnetic and optical methods and it was determined that MGP biofunctionalization with HIV1-TAT yielded the best cellular uptake, from this small CPP screen. The insights provided by this work have potential impacts on the design and optimization of intracellular therapeutic nanoparticle delivery and monitoring for diverse cargos by cell penetration.

5.2 Materials & Methods

5.2.1 Materials

All chemicals used in the synthesis and functionalization of the MGPs were purchased through Sigma-Aldrich unless otherwise noted.

Peptides were purchased through RS Synthesis the sequences are as listed below:

CAAKA: NH$_2$-CAAKA-COOH
Cys-TAT: NH$_2$-C*YGRKKRRQRRR-COOH
His$_6$-TAT-FITC: NH$_2$-HHHHHH*YGRKKRRQRRR(FITC)-COOH
Cys-Ku70: NH$_2$-C*PMLKE-COOH
Cys-hCT(9-32): NH$_2$-C*LGYTYTQDFNKFHTFPQTAIGVGAP-COOH
Cell culture supplies were purchased from Life Technologies, including: RPMI1640 media, Fetal Bovine Serum (FBS), Antibiotics/Antimycotics, Lipofectamine2000®, and T-25 culture flasks. Optical dishes consisted of 10 cm², 1.5 glass bottom, poly-L-lysine coated circular dishes that were purchased from MatTek Corp.

5.2.2 Synthesis of 6nm Fe₃O₄ Cores

Iron oxides nanoparticles were synthesized through the coprecipitation of Fe²⁺ and Fe³⁺ aqueous salt solutions by the supplement of a base, according to modified literature protocols. Briefly, 2 mmol Fe(acac)₃, 10 mmol 1,2-dodecanediol, 6 mmol oleic acid (purity 90%), 6 mmol oleylamine (purity >70%), and 20 mL benzyl ether were stirred under inert N₂ atmosphere. The solution was heated to 200°C for 2 h then, refluxed at 300°C for 1 h. Formation of the Fe₃O₄ nanocores was evidenced by the solution turning black. At which time the nanocores then slowly cooled to room temperature, and precipitated with toluene and methanol. The iron oxide nanoparticles isolated using magnetic separation and resuspended in chloroform for the gold shelling step.

5.2.3 Synthesis of MGP by Gold Shelling of Iron Oxide Cores

A solution consisting of 85 mg of HAuCl₃ and 10 mL of chloroform was prepared by sonication and kept at 4°C for the addition process. The gold salt solution was added to the iron oxide solution at 1 mL per 30 min intervals at room temperature. After complete addition of the gold salt, 100 mg of sodium borohydride was added to the solution to help further reduce the Au on the surface of the Fe₃O₄ core particles to form a contiguous Au shell. After 30 min of sonication, the MGPs were precipitated with toluene and methanol. The MGPs were isolated by magnet separation and dried for the ligand exchange step.

5.2.4 Phase Transfer and Aqueous Bio-Functionalization of MGPs

Approximately 5 mg of MGPs were dissolved in 1 mL of chloroform mixed with 20 mg of thioctic acid (or dihydrolipoic acid: DHLA), deprotonated using tetraethyl ammonium hydroxide and reduced using tris(2-carboxyethyl)phosphine hydrochloride (TCEP), in 1 mL sparged H₂O. The solution was heated to 50°C and stirred for 15 min to allow for phase transfer of the MGP from the organic to aqueous upper layer. Similarly, 2 mg of peptides were dissolved
in 1 mL sparged water and reduced with 50 mM TCEP at 10% volume for 15 min prior to addition to the MGPs. Peptides were appended to the MGP surface by stirring for 30 min to 24 h (variation in timing due to the physical properties of the ligand/peptides, such as charge and MW), with the aide of 300 μL of TEA-OH to deprotonate the carboxylic acids of the peptide and promote better phase transfer. After each reaction, water soluble MGPs were isolated from the top layer and were precipitated with ethanol, followed by centrifugation and magnetic separation.

Dye-labeling of the DHLA and CPP coated MGPs was performed by \(N\)-(3-Dimethylaminopropyl)-\(N'\)-ethylcarbodiimide hydrochloride (EDC) and \(N\)-Hydroxysuccinimide (NHS) coupling of the primary amine containing dye, Nile Blue 690, and the carboxylic acid containing MGP-DHLA, according to the manufacturer’s instructions. All samples were cleaned up by ethanol precipitation and magnetic separation.

5.2.5 MGP Characterization

Synthesis of the Fe\(_3\)O\(_4\) cores and MGPs were characterized by JEM-ARM200cF Transmission Electron Microscope (TEM) for shape, size dispersity and shell uniformity. Crystallinity was confirmed using a Rigaku Ultima3 Powder X-Ray Diffractometer (pXRD). Ligand exchange and aqueous functionalization were verified using Varian Cary 50 Bio Ultraviolet-Visible Spectrophotometer (UV-Vis), Varian Cary Eclipse Spectrophotofluorometer (PL), and a Bruker ALPHA Fourier-Transformed Infrared Spectrometer (FTIR). Magnetic moment and susceptibility were measured by Quantum Design MPMS Superconducting Quantum Interference Device (SQUID) Magnetometer.

5.2.6 Cell Culture and Transfection

A375 skin cancer cells (gifted by Dr. Diego Zorio) and 9L rat gliosarcoma cells (ATCC) were grown at 37°C with 5% CO\(_2\) in RPMI1640 and DMEM7777, respectively, with 10% FBS and streptomycin/penicillin (Gibco). Cells were plated at 10,000 cell/cm\(^2\) into 10 cm\(^2\) optical dishes, or T-25 flasks for cell pellet SQUID measurements, 24 hours prior to transfection. The MGPs were transfected into A375 skin cancer cells by addition of 6.0 pmol/cm\(^2\) (2 mM or 6.7 μg/cm\(^2\)) MGP. A control transfection was performed with the MGP-DHLA using Lipfectamine2000®, as a commercial standard to compare uptake as measured by cellular magnetic susceptibility after transfection versus peptide coated MGP. Lipofectamine2000®
encapsulation of the MGP-DHLA was carried out using a concentration of 150 pmol MGP-DHLA in 10 µL Lipofectamine, per T-25 flask.

5.2.7 Fluorescent Microscopy Cell Imaging

A375 cells were imaged live at 0.5, 2, 4, 8, 12, and 24 h after transfection on a Nikon TE2000 inverted fluorescence microscope. The MGP-DHLA-NB690 was visualized using a Cy5.5 filter cube with exposure time of 900 ms. The MGP-His₆-TAT-FITC and MGP-45FITC were visualized using a FITC filter cube with exposure time of 400 ms. Differential interference contrast (DIC) images were acquired at 100 ms exposure time for visualization of cell boundaries and health. All images were acquired at 40x magnification with a Plan APO short working distance 0.14 numerical aperture objective.

5.2.8 SQUID Measurement of Transfected Cells

MGP transfected cell pellets were prepared for SQUID analysis by collection 24 hours after MGP transfection. Transfection media was removed, cells were washed with TBS three times, prior to treatment with TrypLE (Gibco) for enzymatic cleavage of the adherent cells from the culture dish, with 2 minutes incubation at 37°C. Cells detachment, noted by increased cellular circularity, was visualized under a phase contrast microscope and TrypLE was quenched by addition of a known volume of serum containing medium. Any cells remaining attached were then lifted by gentle rinsing with culture media. Cells were counted and assessed for viability via trypan blue staining using a CEDEX cell counter. Cell solutions of 5 million cells were then centrifuged for 5 minutes at 5K RCF until a dark (MGP-filled) cell pellet was observed. The supernatant was removed the cell pellet was collected and dried in pre-massed high throughput NMR tubes (VWR), for SQUID analysis. Field-dependent studies were recorded at 300 K, with the applied field varying from zero to one Tesla.

5.2.9 MRI Imaging of MGP Uptake in a Tissue Mimic

9L rat glioma cells were transfected 100 pmol of MGP-CPP or MGP-DHLA per well of a 6-well plate. 24 h after transfection, cells were collected resuspended in low melting 2% agarose to create a tissue mimic. Briefly, suspended cells were layered with 150,000 cells per layer with
1% agarose used in between each cell layer, in an NMR tube (VWR). Tissue mimic samples were analyzed using a Bruker MRI at the Nation High Field Magnetic Laboratory.

5.2.10 Optical and Magnetic Analysis

Optical images were analyzed in FIJI ImageJ software\textsuperscript{153} where the total fluorescent intensity was measured in triplicate at each time point and values were corrected for cell number. Curve fitting was performed in the Igor software program to determine uptake kinetics.

Magnetic data was plotted in Microsoft® Excel as magnetic moment per cell versus field strength. Saturation moment was calibrated for each transfection construct and used to calculate relative cellular uptake.

5.3 Results and Discussion

5.3.1 Synthesis of Magnetic Gold Particles

Magnetic gold nanoparticles (MGPs) were engineered by lyothermal synthesis of the Fe\textsubscript{3}O\textsubscript{4} magnetic core with diameter of 6.5 nm, followed by subsequent reduction of HAuCl\textsubscript{4}•3H\textsubscript{2}O salt onto the surface by slow addition in the presence of the reducing agent oleylamine, and finalized by treatment with the strong reducing agent NaBH\textsubscript{4} to encourage a contiguous Au outer shell. Physical characterization of the Fe\textsubscript{3}O\textsubscript{4} cores and Fe\textsubscript{3}O\textsubscript{4}@Au MGPs is shown in Figure 5.2. From the TEM and size distributions (Figure 5.2A-D) it can be seen that the MGP has approximately 0.4 nm of continuous Au shell, indicative of ~1 atomic layer of Au (unit cell: 407.8 pm)\textsuperscript{293} protecting the Fe\textsubscript{3}O\textsubscript{4} core. However, the UV-visible absorption and pXRD data observed display nontrivial indications of an Au shell in the form of the appearance of a plasmon near 520 nm (Figure 5.2E), additional large peaks in the X-ray diffraction matching the Au standard (Figure 5.2F), and reduction of the magnetic moment per sample mass (from 48.1 emu/g to 25.1 emu/g) due to the addition of the heavy and non-magnetic Au shell (Figure 5.2G). This may possibly indicate that the Au reduction onto the Fe\textsubscript{3}O\textsubscript{4} core does not simply add to the net diameter but may, in actuality, etch away layers of the Fe\textsubscript{3}O\textsubscript{4} to allow for a thicker Au shell without greatly increasing the overall NP diameter. Similar atomic exchange phenomenon have been seen previously in other nanomaterial synthesis strategies.\textsuperscript{294}
Figure 5.2 Characterization of the Fe$_3$O$_4$ cores (black) and Fe$_3$O$_4$@Au “Magnetic Gold Particles” (red), showing TEM (A & C) and size distributions (B & D), as well as UV-Vis absorption (E), pXRD patterns (F), and SQUID magnetometry (G). Reference pXRD powder patterns in (F) are for bulk Au (garnet) and Fe$_3$O$_4$ (gray).
5.3.2 Biofunctionalization of Magnetic Gold Particles

Cell penetrating peptides (CPP) have been observed to facilitate cellular entry for a variety of nano- and molecular-cargos.\textsuperscript{90} As mentioned in the previous chapter, the presentation and packing of the CPPs on a the NP surface has been proposed as an explanation for the observed selectivity of CPPs for various cell lines. Selection of known CPP sequences from the literature such as the human calcitonin peptide hormone (hCT), the transduction fragment of the Bax inhibiting protein Ku70, and, the transactivator of transcription for the human immunodeficiency virus-1 (HIV1-TAT) were chosen to provide a small representative screen of large (hCT: 24 residues, +1 charge), small (Ku70: 5 residues, +1 charge), and highly charged (TAT: 11 residues, +6 charge) CPPs known to induce cellular internalization. In addition, the sequences were mutated to contain N-terminal cysteine modifications (denoted as C*) to facilitate thiol binding to the Au surface of the MGP, and co-loaded at a 1:9 ratio with the small synthetic peptide, CAAKA, in order to optimize packing and sequence presentation on the MGP surface. Additional appendage chemistries exist including the amine-based hexa-Histidine (H\textsubscript{6}) coordination, which allows for six consecutive N-terminal histidine residues to bind to the surface of thee inorganic NP, though dative bond formation with the imidazole rings of the histidine.

In Figure 5.3A, it can be seen that while short peptides (CAAKA and C*Ku70) are not likely to generate notable secondary structure on the MGP surface, the most energetically favorable orientations of the larger peptides (C*hCT and C*TAT) entail formation of some degree of secondary structure with an overall reduction in the total “height” of the CPP from the MGP surface, as compared to the primary amino acid sequence, which may play a role in cell surface binding and ultimately in cellular internalization efficiency. As a comparator and small molecule drug mimic, dihydrolipoic acid (DHLA) – the reduced form of the vitamin α-lipoic acid, known to mitigate oxidative cellular stress\textsuperscript{295} – was also used in this study (Figure 5.3A). The addition of the biologically useful CPPs and the small-molecule drug-mimic to the surface of the MGP was accomplished by biphasic ligand exchange, where the as-synthesized organically soluble MGPs were dispersed in CHCl\textsubscript{3} and layered with a basic (pH 10) aqueous solution of reduced cysteine-modified (C*) CPPs or dihydrolipoic acid (DHLA) to provide reactive thiols for binding the Au surface coating of the MGP (Figure 5.3A & B).
Figure 5.3 Ligand exchange characterization of MGPs with various cell penetrating peptides and the small molecule DHLA (A) showing structures; MGP (B-i) organic vs. (B-ii) aqueous solubility, before and after ligand exchange, respectively; and (C) absorption profiles for water soluble biofunctionalized-MGPs.
The reactions proceeded rapidly at 50°C, between 10 and 60 min, with an apparent positive correlation between the time to completion and the molecular weight, compounded by charge, of the appending aqueous molecule (t_{rxn}: DHLA<CAAKA<C*Ku-70<C*hCT<C*TAT).

Notably, place exchange reactions allowed to proceed longer than 8 hours yielded aggregated MGPs, most evident with the highly charged C*TAT peptide (+6). This may be due to possible weakening of the Au shell from bombardment with reactive thiols over long time spans, allowing for fusing of the unprotected magnetic cores, or potentially due to peptide-peptide interactions inducing conglomeration of the particles. However, the shorter reaction times, yielded water-soluble MGPs with visible surface plasmon resonance near 525 nm (Figure 5.3C). For the shorter peptide functionalization (CAAKA and C*Ku70) the plasmon is not as pronounced as for MGPs with the larger peptides (C*hCT and C*TAT) as well as for the DHLA, likely due to differences in the local electronic environment induced by the differences in peptide and small molecule electronegativity and hydrophobicity. Additionally, some broad absorbance due to possible MGP agglomeration is evident for the highly charged C*TAT and the large C*hCT in the ~600 nm region, not evident in the MGP-CAAKA, MGP-C*Ku70 and MGP-DHLA samples, likely due to bio-ligand size and charge.

5.3.3 Magnetic Resonance Imaging of MGP-CPPs in a Tissue Mimic for Gliosarcoma

The ability to assess uptake and localization of therapeutic materials, to determine therapeutic efficacy, in clinical settings can present a challenge for certain cancers, such as glioblastoma, which are not easily visibly accessible without invasive procedures. However, the magnetic core of the MGP allows for visualization via Magnetic Resonance Imaging (MRI), as the MGP acts as a magnetic contrast agent versus the mostly water-filled cell and tissue volume. Here, CPP and DHLA coated MGPs were used as therapeutic analogs, transfected into cultured 9L rat gliosarcoma cells, and were able to be imaged via MRI in a tissue mimic composed of cell layers suspended in an agarose gel matrix (Figure 5.4). The CPP loaded MGPs were transfected without the aid of any commercial transfecting agent, but the MGP-DHLA, which is unable to act as a membrane penetrant, was encapsulated using Lipofectamine2000® (denoted as MGP-DHLA_{lipo}) to facilitate cellular entry into the 9L cells. The quantification of cellular uptake for the MGP constructs was performed by assessing the relative relaxation rates for the MGP treated cells versus the agarose background. The T1 recovery relaxation time is a
measure of the ability of the sample to recover its spin orientation with the applied magnetic field after being exposed to a transverse magnetization pulse of radio frequency radiation. Figure 5.3A shows that all of the MGP-CPP and MGP-DHLa<sub>lipo</sub> samples display indistinguishable T1 relaxations from the agarose control. T1 relaxation describes the rate of the redistributions of the nuclear spin state populations as they become representative of the thermal equilibrium, of which the signal here is dominated by the water background. The T2 decay relaxation, however, is a measure of the amount of time needed for the effects of the transverse magnetization radio pulse to dissipate, or how susceptible the sample is to “forgetting” the transverse magnetization pulse in the presence of the applied field, which will inherently be shorter for a magnetic material.

![Graphs and images from Figure 5.4](image_url)

**Figure 5.4** Magnetic resonance imaging analysis of cellular uptake for MGP-CPPs and MGP-DHLA (transfected using Lipofectamine2000®) constructs, prepared in agarose gel tissue mimic. Quantification and imaging of T1 Relaxation (A & D), T2 Relaxation (B & E), and T2* Relaxation (C & F), as well as 3-D model of MGP localization in the tissue mimic (G).
Here, stronger differences are seen between the T2 of the MGPs and the agarose (Figure 5.4B), and upon inspection of the T2* decay relaxation, which takes into account inhomogeneities of the magnetic field, it can be seen that the shorter relaxation times of MGP-C*TAT and MGP-DHLA\textsubscript{lip} indicate a higher degree of magnetization (i.e. a larger number MGPs) compared to the other samples, and that all of the MGP samples are significantly different from the control agarose (Figure 5.4C). Correlating MRI visual inspection of the MGP-CPPs and MGP-DHLA\textsubscript{lip} cellular uptake in the agarose tissue mimic, T1, T2 and T2* weighted images are observed to corroborate the computed uptake data, with the MGP-C*TAT and MGP-DHLA displaying the darkest contrast in the T2 and T2* weighted images (Figure 5.4E & F), and the T1 image showing no strong differences between samples (Figure 5.4C). The three-dimensional MRI image (Figure 5.4G) displays granulated areas of high contrast for all of the MGP samples, indicating possible clustering of transfected cells due to magnetic-attraction of the MGP-containing cells during the process of suspending in agarose, following the 24 h incubation with the MGP samples.

5.3.4 Optical Functionalization of the MGPs with FITC and Nile Blue690

The multi-functionality of the MGP as a magnetic contrast agent in the difficult-to-visualize deep-tissue gliosarcoma tumor mimic led to the finding that the MGP-C*TAT and MGP-DHLA in Lipofectamine\textregistered displayed the highest intracellular concentrations of magnetic particles. However, since the resolution of MRI does not allow for individual cell-level information to be obtained, investigation of the utility of the plasmonic Au shell of the multi-functional MGPs was conducted. A375 human skin cancer cells were cultured as a model of an accessible cancer type, melanoma, where a biopsy could be easily collected and cultured to assess MGP therapeutic efficacy using fluorescence microscopy. When electing for the use fluorophores to functionalize a MGP for fluorescent analysis applications, one must consider the plasmonic Surface Energy Transfer (SET) quenching behavior of solid AuNPs of similar size.\textsuperscript{130} Since the 6 nm MGP will have similar quenching behavior to a solid 6 nm diameter AuNP, fluorophores blue of the plasmon at 525 nm will experience quenching dependent on their emission wavelength and distance from the MGP surface. Conversely, fluorophores red of the 525 nm plasmon will not engage in SET coupling and will remain fully “on.”\textsuperscript{52,130}
Figure 5.5 Optical functionalization of best performers, MGP-TAT and MGP-DHLA with fluorescein isothiocyanate (FITC) and Nile Blue690 (NB), respectively. Showing absorbance of (A) MGPs before (OA/OA) and after biofunctionalization with DHLA, C*TAT and H6*TAT, as well as (B) before and after coupling NB to DHLA. Additionally, for the SET quenched FITC, treatment with sodium cyanide to etch the Au shell show changes the (C) absorbance and (D) photoluminescence profiles after 5 minutes and full Au dissolution and loss of the Au plasmon as well as FITC fluoresce recovery by 2h.

For this reason, the two fluorescent dyes selected to functionalize the MGPs were fluorescein isothiocyanate (FITC: $\lambda_{\text{EXC}}$ 495 nm, $\lambda_{\text{EM}}$ 519 nm) and Nile Blue690 (NB: $\lambda_{\text{EXC}}$ 635 nm, $\lambda_{\text{EM}}$ 674 nm). For the fluorescent microscopy studies, the best performers, TAT and DHLA, were fluorescently labeled. FITC labeled TAT, mutated with an N-terminal hexa-histidine sequence (H6*TAT-FITC), was purchased due to the less stable nature of imidazole coordination with the Au shell, from the histidine residues, as compared to the strong thiol-Au bond formed by the N-terminal cysteine used previously. The stability of the peptide-MGP bond becomes important for the decoupling of the SET quenching of the FITC, as a strong peptide-MGP bond will not be conducive to peptide release from the MGP surface, and thus fluorescence recovery and
subsequent detection would not be possible. The NB was appended to the free carboxylic acid of the DHLA using EDC coupling chemistry. Due to the short distance of the DHLA as a spacer from the Au shell, and the strength of dual thiol-Au bonds, a red dye was used to ensure detectability. Figure 5.5 displays the absorbance characterization data for the MGP-H₆*TAT-FITC (A) and MGP-DHLA-NB (B), comparing MGPs before and after dye labeling. The absorbance profiles of the FITC and NB dyes can be seen in the final coupled products (Figure 5.5A & B). Additionally, since the FITC is quenched by the Au shell of the MGP, the use of cyanide to dissolve the Au shell was employed to test fluorescence recovery of the FITC after binding to the MGP, where both the absorbance spectra display a loss of the Au shell plasmon, and the emission profiles display a gain in FITC intensity within 5 minutes, and full recovery after 2 h (Figure 5.5C & D).

After successful dye-functionalization, the labeled MGPs were incubated with A375 melanoma cells for 24 h. Live cell fluorescence microscopy allowed for the MGP-H₆*TAT-FITC and MGP-DHLA-NB transfection progressions to be imaged over the 24 h experiment. Figure 5.6 depicts the health of the cells following transfection via the differential interference contrast (DIC) images (gray), as well as the change fluorescence intensity and cellular localization of the FITC (green) and NB (red), in the fluorescence and overlay channels, respectively (Figure 5.6). As expected from the predicted quenching behavior of the Au shell, the FITC shows a gradual increase in fluorescence intensity over time, as H₆*TAT-FITC is released from the MGP surface, reaching full intensity at 8 to 12 h post transfection (Figure 5.6A). Alternately, the NB fluorescence intensity is observed to be fully “on” at the earliest time point, as can be seen in the 2 h images (Figure 5.6B). Also worth noting, the fluorescence pattern of the TAT facilitated MGP uptake displays a more punctate pattern at early time points, indicative of endosomal entrapment, followed by whole cell diffuse fluorescence at later time points, indicating cytosolic escape of the fluorescent cargo. Whereas, the MGP-DHLA in Lipofectamine2000 shows a higher degree of cytosolic localization at earlier time points, possibly pointing to an advantageous ability of DHLA modified particles to avoid or escape endosomal entrapment even when delivered in a cationic liposome. Extraction of the fluorescence intensity values for the FITC and NB channels over time yields the uptake plot shown in Figure 5.7. Here it can be seen that the NB (red) intensity is strongly observed from the earliest time points, with the slight delay in full
Figure 5.6 Fluorescence microscopy images of A375 melanoma cells at 2, 4, 8, 12, and 24 h post-transfection with (A) MGP-TAT-FITC and (B) MGP-DHLA-NB. Differential interference contrast (DIC), fluorescence, and composite overlay images are shown for each time point. Scalebar: 100 μm.

emission being likely due to the settling of liposomes onto cells and the time needed for cellular internalization, to enter the focal plane under observation (Figure 5.7). Additionally, the NB maintains a relatively constant emission profile throughout the experiment, with only slight decreases in intensity per cell occurring due to changes in cell growth over time and no additional supplementation of MGP-DHLA-NB (Figure 5.7). The FITC intensity, on the other hand, is observed to follow a much more gradual sigmoidal turn-on, which reaches its maximal value at 8 h post transfection, and remains constant despite changes in cell growth, indicating potential continued slow release of the H₆*TAT-FITC from the MGP (Figure 5.6). Concurrently, the fluorescence turn-on half-life for the MGP-DHLA-NB was found to be ~30 min, as compared to the half-life of FITC which was found to be ~3 h. Previous work with DNA shows similar cargo release kinetics from a solid AuNP.⁵⁶
Figure 5.7 Analysis of A375 cellular internalization of the MGP-bioconjugates via microscope fluorescence for MGP-DHLA-NB (red ○) and MGP-H₆*TAT-FITC (green ●) as total intensity per cell, over the 24 h experiment. Corresponding sigmoidal fits indicated by solid lines for the MGP-DHLA-NB (red) and MGP-H₆*TAT-FITC (green) data points, respectively.

Surprisingly, the overall intensity of the NB fluorescence is observed to be much higher than the FITC, despite large counteracting differences in quantum efficiencies (FITC 92% vs. NB 27%) and detector sensitivities (CoolSnap HQ2 CCD: 519 nm ~55-60% vs. 674 nm ~25-45%) for each dye. While a decreased intracellular pH being experienced by the TAT-FITC, as in the cases of lysosomal sequestration, may be responsible for the reduced FITC intensity, a more likely potential cause may be owed to differences in labeling efficiency and ligand loading levels on the MGP-H₆*TAT-FITC versus the MGP-DHLA-NB. The former is loaded at 1:9 with the unlabeled CAAKA peptide, versus the latter which is passivated by the smaller molecule, DHLA, which coats the entire MGP surface. The attachment of the NB dye by the method of EDC coupling is known to be moderately inefficient, however, the numerical difference of available addition sites for the MGP-DHLA is likely overwhelming when compared to the labeling of the MGP-H₆*TAT-FITC. Alternatively, the observed lower FITC signal could be indicative that not all of the FITC labeled H₆*TAT has been released from the MGP even by the 24h time point. Thus, the optical data provided by the utility of the plasmonic Au shell of the
MGP can provide insights on the localization and kinetics of intracellular delivery, but may not be able to accurately quantify cellular uptake due to potential convolution with differential labeling or incomplete cargo release.

5.3.5 MGP Cellular Uptake Assessed by Magnetometry

In an attempt to better assess and more accurately quantify the number of MGP taken up by the A375 melanoma cells, for the TAT versus DHLA bio-functionalization approaches, which gave the best performance in the MRI study, analysis of isolated cell pellets was performed 24 h after transfection using a Superconducting Quantum Interference Device to measure magnetic moment saturation compared to the number of cells. In Figure 5.8 it can be seen that the MGP-C*TAT shows more than twice the magnetic moment as compared to the MGP-DHLA\textsubscript{lipopo} for a sample of five million cells, respectively. This gives an approximate value of 3.40x10\textsuperscript{8} emu/cell for the MGP-C*TAT as compared to 1.48x10\textsuperscript{8} emu/cell for the MGP-DHLA\textsubscript{lipopo}, which signifies an increase of 2.3:1 in MGP uptake for the CPP loaded versus the small molecule loaded MGP (transfected using lipofection) (Figure 5.8).

![Figure 5.8](image)

**Figure 5.8** Superconducting Quantum Interference Device (SQUID) measurements of magnetic moment saturation for collected A375 cell pellets (5 million cells) at 24 h post transfection with MGP-C*TAT (♦) and MGP-DHLA\textsubscript{lipopo} (□). The “M” on the y-axis represents magnetic moment and the “H” on the x-axis represents magnetic field strength.
Comparison to the calculated magnetic moment per particle (~5.9x10^{-17} emu/MGP), as corrected for approximately 45% ligand mass and assumed to be relatively unaffected by particle size distributions, led to the finding that the MGP-C*TAT experienced an average cellular uptake of approximately 580 million particles per cell, whereas the MGP-DHLA_{lipo} facilitates internalization of 250 million particles per cell. This demonstrates the efficiency of MGP uptake to be greater when using the CPP TAT versus the commercially available cationic lipid transfection methodology, with approximately 80.3% of transfected particles entering cells for MGP-C*TAT, and approximately 35.0% of MGP-DHLA entering cells with the assistance of the lipofectamine transfection reagent.

Analysis of the other MGP-CPP constructions gave undetectable magnetic moment signal over the background when assessed with the large mass of the cell pellets. Differences in CPP-dependent uptake patterns from previous work on semiconducting quantum dots may be due to changes in the achieved loading level or CPP packing on the Au shell versus the ZnS shell of the QD, emphasizing the need for individual material analyses.

5.4 Conclusion

In summary, the synthesis and bioconjugation of multi-function magnetic gold nanoparticles from Fe$_3$O$_4$ cores coated with a Au shell, followed by appendage of modified CPPs and DHLA, as a small molecule drug mimic, has been demonstrated. Investigation of MGP-CPP and MGP-DHLA_{lipo} uptake in 9L rat gliosarcoma was assess in a layered agarose tissue mimic by MRI; and it was found that the MGP-C*TAT and MGP-DHLA (transfected by lipofection) showed the highest T2 and T2* contrast, indicating greater cellular uptake versus the CAAKA, Ku70 and hCT peptides. Utilization of the plasmonic Au shell of the MGP allowed for assessment of the cellular localization and uptake kinetics for the best performers of MGP-TAT and MGP-DHLA in a melanoma biopsy model of A375 skin cancer cells. More rapid and intense fluorescence was observed by the MGP-DHLA-NB_{lipo} as compared to the MGP-H$_6$*TAT-FITC which had a half life of ~3 h versus 0.5 h for NB. This difference maybe partially due to the “always on” emission of the NB, since it is red of the Au plasmon and not energetically coupled by SET quenching, versus the SET quenched FITC appended to the MGP-H$_6$*TAT which required release of the coordinated imidazoles of the hexa-histidine anchoring group, prior to FITC emission recovery. Further quantification of MGP uptake was assess via SQUID
measurements of magnetic moment saturation per cell which found that the MGP-C*TAT outcompeted the MGP-DHLA_lipo for cellular uptake in A375 cells by 2.3:1 with 80.3% and 35.0% uptake efficacy, respectively.

The findings of this work not only establish MGPs as a multi-utility bio-functional platform for cellular transfection, treatment, and diagnostic applications, but also demonstrate multiple magnetic- and plasmonic-analysis approaches for probing clinically relevant nano-therapeutic questions of intracellular fate and internalization efficacy. In all, the bio-conjugated MGP has been shown to have great potential as a clinically-valuable nanotechnology.
CHAPTER 6
SUMMARY AND FUTURE OUTLOOK

The body of this work described the design, fabrication, implementation, and analysis of several unique applications for nanomaterials as biological sensors in living cells, used as models for nanoparticle-based therapeutic approaches. Utilizing the molecular beacon toolbelt, the nano-specific optical and magnetic properties, and the benefits of peptide-mediated cellular targeting and entry, spectroscopic and microscopic analysis of cellular uptake, processing, and fate was accomplished. The fundamental science explored herein helped to elucidate important factors involved in nanomaterials-bioconjugation as well as those pertaining to intracellular processing of nano-probes and nano-therapeutics.

In chapter two, the study of gold nanoparticles (AuNPs) used as delivery platforms for DNA oligonucleotides and genes with various appendage strategies revealed the ability to engineer control over the attachment chemistry and thus the release rate of DNA therapeutic cargos. Here the common mono-thiol oligonucleotide-AuNP conjugation strategy was compared to dual thiol, dual amine, and thiol plus amine bidentate conjugation schemes as well as mono-amine conjugation and it was determined that differential rates of intracellular processing and therapeutic release could be accomplished for each coupling method. The use of live-cell fluorescence microscopy allowed for the direct visualization of the timing of release through fluorescence “turn on” of the dye-labels on the DNA that were quenched by the AuNP while the DNA was bound to the particle within the nanometal’s Surface Energy Transfer (SET) diameter. The intracellular rates of DNA release from the AuNP surface in A375 human melanoma cells were found to follow the binding activity series: dual thiol > thiol plus amine > mono thiol > dual amine ≈ mono amine, for the investigated DNA- AuNP coupling routes. And similar trends were observed for gene expression of a linearized plasmid containing a fluorescent protein (FP) gene appended by the same strategies. Future endeavors to expand this work will be able to implement active therapeutics such as a gene for a short hairpin RNA which will continuously knock-down expression of a delinquent protein, such as the P-glycoprotein efflux pump which is known to be upregulated in multi-drug resistant cancers, followed by a secondary release event to deliver small molecule drug, or suicide gene to terminate the treated
cancer cell, from the same AuNP. These experiments could also be assessed by live-cell fluorescent microscopy using a FP gene fusion, or via other cell viability assays.

In chapter three, the simultaneous monitoring of the timing of cargo release from the nanotherapeutic AuNP as well as measurement of the evolving intracellular pH experienced by the cargo during cell processing was elucidated. This was accomplished using a triple dye-labeled duplex DNA appended to the AuNP that employed the environmentally sensitive and pH-responsive dye, fluorescein, paired with DyLight405, to act as ratiometers in measuring intracellular pH, at and after cargo release; as well as DyLight700, which was not energetically-coupled to the AuNP, as a measure of uptake and localization throughout the experiment. Through visualization via live-cell fluorescent microscopy, it was observed that the nanotherapeutic packaging of labeled DNA provides a protective effect against lysosomal degradation in acidic organelles as noted by cargo experiencing an average intracellular pH near 6. Future expansion of this sensor platform may encompass multiple spectrally resolved sensing dyes, appended to as single AuNP, to multiplex intracellular analyte detection and better elucidate the complete picture of the cellular environment experienced by the nanoparticle, including the presence and abundance of reactive oxygen species, and reducing agents encountered by the nano-therapeutic.

In chapter four, the role of drug resistance status in cancer cells was probed by the targeted therapeutic delivery of emissive quantum dots (QDs) biofunctionalized with cell penetrating peptide (CPP). The QD-CPP nano-bioconjugation created a labelled delivery package to investigate the nuances and difficulties of drug transport into multidrug resistant (MDR) cancer cells for potential future clinical applications of diverse nanoparticle therapeutic delivery strategies. The eight distinct CPPs used were CAAKA, VP-22, HIV-TAT, HIV-gp41, Ku-70, and hCT(9-32), integrin-β3, and K-FGF. Differential cellular uptake profiles were observed in cancer versus drug resistant melanoma (A375 & A375-R), mesothelioma (MSTO & MSTO-R), and glioma (rat 9L & 9L-R, and human U87 & LN18), cell lines, with hCT being the best performer for most cell lines, and showing stronger affinity in drug-resistant melanoma and gliosarcoma. The results also demonstrated that cell penetrating peptide uptake varies with the amount of drug resistance and cell type, likely due to changes in cell surface markers. This study provided insight into developing functional nanoplatform delivery systems in drug resistant cancer models. Future expansion of this sensing platform will include delivery of
chemotherapeutic and anticancer genes to assess how uptake and release rates for the nano-
therapeutic payload are affected by co-assembly with the CPPs.

In chapter five, the collection of analytical methodologies available for evaluating nano-
therapy in cells transfected with gold shelled magnetic nanomaterials, termed “magnetic gold
particle” (MGP), was demonstrated. Analysis of the MGPs’ magnetic iron oxide core, enabled
evaluation of magnetic moment and resonance via SQUID and MRI, respectively, to determine
MGP-CPP uptake levels in a deep tissue MRI model, as well as magnetometry of an extracted
cell pellet to allow multipoint analysis of NP internalization. The added advantages of a gold
shell allowed for good biocompatibility, easy biofunctionalization with the CPPs, and
visualization of intracellular therapeutic delivery and kinetics using fluorescent microscopy for
optically functionalized MGPs with fluorophores possessing SET and non-SET behavior. This
multiplex sensor demonstrated the best aspects of magnetic and gold nanoparticles in one useful
platform, and addressed questions of therapeutic targeting and uptake with cell penetrating
peptides in clinically relevant cancer cell models. Future exploration of the full utility of the
MGP may include implementation into in vivo clinical models, such as brain and skin cancer
tumors, expressed in mice, as well as employment of functional therapeutics to co-append with
the CPPs to drive therapeutic benefit of the MGP platform to the next level.

In all, this work has set an important foundation for the future, and present, clinical
development of nanotherapeutics for personalized cellular medicinal approaches. Understanding
how, where, and why nanoparticle-bioconjugate constructions are processed by individual cells
is paramount to the success of nanomedicine. Being aware of potential perils, such as lysosomal
degradation of the nano-cargo, as well as being able to utilize advantageous outcomes of
intracellular processing of nanomaterials, such as control over cargo-release and cell targeting,
will drive better design and multifunctionality to be engineered into the next evolution of
medicine.
APPENDIX A

ABBREVIATIONS AND ACRONYMS

AuNP, gold nanoparticle
BOT, Bio-optical Transponder
CCF, cross-correlation function
DNA, deoxyribonucleic acid
DL, dylight
EE, early endosome
FAM, fluorescein amidite
LE, late endosome
LY, lysosome
M1, Manders’ colocalization coefficient.
McSET, multicolor Surface Energy Transfer
NP, nanoparticle
NA, nucleic acid
r_p, Pearson’s correlation coefficient
RFP, red fluorescent protein
SET, Surface Energy Transfer
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BIOGRAPHICAL SKETCH

Katelyn Jo Fredrickson [Carnevale] was born in Twenty-nine Palms, CA in 1988 and attended high school in Jacksonville, FL from 2002-2006, graduating in the top 3 of her class from Seacoast Christian Academy, with a 4.1 GPA due to weighted dual enrollment courses taken at Florida State College at Jacksonville (FSCJ). Kate got her Associates degree from FSCJ in 2009, where she was presented with the “Outstanding Student in Chemistry” Award. Kate went on to earn dual Bachelor degrees in Biology and Chemistry in 2012 from the University of North Florida (UNF), in Jacksonville, FL, where she received her second “Outstanding Student in Chemistry” Award. During her undergraduate career, Kate was worked as a Certified Pharmacy Technician from 2008-2012. Kate also conducted undergraduate research in the Chemistry Department at the UNF with Prof. Amy Lane, assaying and isolating novel antibiotic compounds from marine fungi samples.

In August of 2012, Kate enrolled at Florida State University (FSU) in the Department of Chemistry & Biochemistry in the Biochemistry Graduate program and joined the laboratory of Prof. Geoffrey F. Strouse. She began working in the field of nano-biomedicine under the mentorship of Megan Muroski (now Ph.D.). During her time in the Strouse research group, Kate worked at synthesizing and bio-functionalizing nanomaterials for intracellular uptake and gene delivery studies, and assessed nanomaterial behaviors within mammalian cells via optical fluorescence microscopy imaging of emissive and SET nanomaterials, as well as quantification of uptake of magnetic nanomaterials via MRI and SQUID analysis. She was awarded the Joseph M. Schor Fellowship in Biochemistry in 2015 in recognition of her graduate work. Kate has also presented her work at local, regional, and national meetings at: the FSU Life Sciences Research Symposia (2013 & 2015 in Tallahassee, FL), the FSU Natural Sciences Graduate Research Symposium (2014 in Tallahassee, FL), the Southeastern Regional Meeting of the American Chemical Society (2015 in Memphis, TN & 2016 in Columbia, SC), and the National Meeting of the American Chemical Society (2015 in Boston, MA & 2018 in New Orleans, LA). And her work has been published in the Journal ACS Nano (2015 and 2018).

Kate has been involved in her scientific community as is a current member of the American Chemical Society (ACS) since 2011, and the Association of Women in Science (AWIS) since 2018. Kate was also the Founder and President of the FSU ChemiNole Graduate Student Association (CGSA) from 2016-2017, and President of the FSU Graduate Women in
Science Student Organization (GWIS) from 2015-2016, as well as President of the UNF Student Chapter of American Chemical Society (ACS) from 2011-2012.

Kate has also served the Department of Chemistry & Biochemistry as the Chair of the Graduate Student Recruitment Committee (2015-2017) and Department Representative for the FSU Program for Instructional Excellence (PIE) (2015-2016), along with her many years as a Graduate Teaching Assistant for General Chemistry, Organic Chemistry, and Biochemistry (2012-2018).

Kate is also very active in her faith community as a Sunday School Teacher, Children’s Ministry Facilitator, and Adult Small Group Leader (2008-2018). And has participated in events to benefit several local charities that serve single mothers (Making Miracles Group Home), and low-income families (Habitat for Humanity), as well as abandoned and abused animals (Jacksonville No More Homeless Pets). Kate also supports several national and international charities that serve the needs of orphans and children in poverty (Convoy of Hope, Stella’s House, One Child Matters, and Compassion International).