The Design of a Molecular Assembly Line Based on Biological Molecules

by

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B.S. Chemistry
Stanford University (2001)

Submitted to the Program in Media Arts and Sciences,
School of Architecture and Planning
in partial fulfillment of the requirements for the degree of
Master of Science in Media Arts and Sciences
at the
Massachusetts Institute of Technology
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Abstract

A general scheme towards a “molecular assembly line” based on biological molecules is proposed, as well as its potential uses as a universal polymer scaffold in programmed assembly and molecular electronics. It is based on the principles of the biological molecules polyketide synthase and kinesin, and in some embodiments, may employ biomolecules like DNA as components of the system. The scheme entails the construction of a polymeric chain that moves a shuttle along the chain by controlling the interactions between the shuttle and individual monomer units using external inputs. The experimental work here particularly focuses on the design and synthesis of the monomer units and shuttles, as well as the mechanisms of control over the monomer/shuttle interactions that are required to construct the proposed polymer systems. Two approaches are explored, the first of which utilizes radio-frequency magnetic fields to selectively dehybridize DNA by coupling RF to covalently attached nanoparticle antennae. The second approach utilizes wavelength selective photocleavage of carbonyl bonds to control the equilibrium of a Michael reaction, and will demonstrate how one can construct a purely synthetic analogue of a polyketide synthase.

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“A chemist comes to us and says, ‘Look, I want a molecule that has the atoms arranged thus and so; make me that molecule.’ The chemist does a mysterious thing when he wants to make a molecule. He sees that it has got that ring, so he mixes this and that, and he shakes it, and he fiddles around. And, at the end of a difficult process, he usually does succeed in synthesizing what he wants. By the time I get my devices working, so that we can do it by physics, he will have figured out how to synthesize absolutely anything, so that this will really be useless.”

- Richard P. Feynman, “There’s Plenty of Room at the Bottom” (1959)
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Chapter 1: Introduction

1.0 – Overview

This thesis discusses the general design of a “molecular assembly line” based on biological molecules for the purpose of programmed assembly at the molecular level. The molecular assembly line is a linear polymer chain that is a scaffold capable of passing a solid support for chemical synthesis along the scaffold by any arbitrary pathway. The scheme is based on biological molecules in that it is based on the principles of biology’s robust assembly processes, and also in that various embodiments of the assembly line may employ biological molecules as building blocks. In addition to outlining a general scheme, this thesis will also present experimental results on some basic building blocks of different types of molecular assembly lines.

Why would one want to create a molecular assembly line? The notion that thousands of “tiny hands” could work in parallel to construct devices by maneuvering individual atoms was first proposed over 40 years ago by Richard Feynman in his prophetic speech - “There’s Plenty of Room at the Bottom” [Feynman59]. The dream was realized in 1993, when IBM researchers [Eigler93] arranged xenon atoms into the IBM logo by using a scanning tunneling microscope (STM) tip to pick-and-place individual xenon atoms on a nickel surface (figure1.1.). After this tour de force, the notion that humans could build complex systems atom by atom no longer seemed like a pipe dream.

“What would the properties of materials be if we could really arrange the atoms the way we want them? They would be very interesting to investigate theoretically. I can't see exactly what would happen, but I can hardly doubt that when we have some control of the arrangement of things on a small scale we will get an enormously greater range of possible properties that substances can have, and of different things that we can do.”

- Richard P. Feynman, (1959)

In recent years, largely driven by the desire to construct physical systems with 2-D assembly and 3-D folding rules like proteins, researchers have been working towards
programmed assembly [Whitesides02], in which various structures can be created from a universal parts set. The notion of programmable chemistry [Lehn00][Stoddart02], the synthesis of various chemicals from universal precursors, has also become popular recently. In these programmable systems, the outcome of the assembly or synthesis depends on the inputs into the system, usually chemical, electrical, or optical. Molecular electronics have also come to prominence in recent years for the vast potential device density and higher performance in devices utilizing quantum size effects. Indeed, if an individual molecule could be used as a switch, one liter of a 1M solution could potentially hold on the order of $10^{23}$ switches, or roughly $10^{11}$ Pentium IV chips, which contain about one trillion elements.

The proposed molecular assembly line will be important to these fields because it represents a system that is truly molecular, scaleable, and programmable. As will be discussed in section 1.1, even though many molecular devices have been demonstrated, these devices often contain only one component that is a molecule, and the rest is lithographically defined on a 2-D surface; the device is more “molecule-based” than “truly molecular” in nature. Truly molecular devices have been shown, but it has yet to be shown how these systems can be built-up. The molecular assembly line will demonstrate how to construct analogs to scaleable and robust biological systems, and it will shed light on how biological molecules can be used for general computational purposes, like the construction of bi-stable molecular switches.

In this chapter, section 1.1 overviews the current state of molecular devices and some of the issues surrounding devices that have been reported in the literature. Section 1.2 describes the biological inspirations for the proposed molecular assembly line. Section 1.3 proposes the general scheme of the molecular assembly line and various embodiments of the system.

Chapter 2 proposes a molecular assembly line based on “RF Biology” – the control of DNA hybridization states by coupling radio-frequency magnetic fields into covalently attached superparamagnetic nanoparticle antennae. Chapter 3 proposes an alternative mechanism of controlling biological molecules using static magnetic fields, termed “DC Biology.” Chapter 4 proposes a molecular assembly line based on wavelength selective
photocleavage of carbonyl compounds. Finally, chapter 5 summarizes the work presented.
1.1 – Issues in molecular devices

In recent years, nanotechnology research has come to significant prominence, but the term is broad and ill defined. Likewise, many terms used in discussing nanotechnology are also ill defined, terms as simple as “molecular.” For example, is a Pentium IV, which currently has a design rule of 110nm, considered “nanotechnology?” Before discussing the merits of and problems with molecular devices, certain arbitrary definitions or requirements must be made to limit the scope of this introduction and thesis.

*The size benefits of devices in nanotechnology* - will be limited to device density. Ballistic transport, electron tunneling, quantum entanglement, and other physical phenomena associated with the nanometer size scale will not be considered, despite their importance in nanotechnology with respect to computational speed, efficiency, energy consumption, and cryptography.

*For a device to be considered “molecular”* - the device must be built by a bottom-up approach; the device cannot be created in its entirety by lithography or any other top-down approach, even if every component is in the nanometer size scale. This rules out Pentiums and other purely lithographically defined devices that have design rules in the nanometer regime.

*For a device to be considered “truly molecular”* – the entire device must be soluble in some solvent after fabrication. The presence of a 2-D surface or substrate must not be absolutely necessary to its performance. This rules out a process like manipulating atoms with a STM tip as shown in figure 1.1.

*The term “programmable”* - entails the ability to affect the system such that there is only one outcome despite there being many potential outcomes initially.

*The term “externally programmable”* - requires a remote input that does not require spatial confinement to address the system elements. Once again, this rules out a Pentium chip that requires wire bonding between lithographically defined electrode structures, or a process like manipulating atoms with a STM tip.
How “scaleable” a device is – is a matter of how large or complex of a system can be built by only adding individual devices to the system being constructed. For example, the ability to construct logic gates out of switching elements is the measure of how scaleable the switch is, but the ability to build processors out of the logic gates does not reflect on the scaleability of the switches. This limitation must be made because the connecting of many higher-order devices does require spatial confinement in almost all schemes proposed in molecular electronics, including externally programmable systems.

It would be impossible to summarize and discuss all molecular machines and devices that have ever been constructed, but here they are put in two categories, molecule-based devices employing lithography, or unscaleable molecular systems in solution.

1.1.1 – Molecule-based devices employing lithography

In recent years, many researchers have been using nanotubes, nanowires, nanoparticles, polymers, and individual molecules to build nanoscale devices like field-effect transistors [Avouris01] by replacing the lithographically defined semi-conductor gap with semi-conducting molecular components (figure 1.2). While there are exciting phenomena associated with this size scale, in particular, tunneling [McEuen96], ballistic transport [Dai01], and quantum entanglement [Herz02], nearly zero Kelvin (liquid helium) temperatures are typically required for these devices to utilize these phenomena.

![Figure 1.2](image.png)

**Figure 1.2** Not-gate (inverter) composed of a semi-conducting single-walled carbon nanotube bridging two sets of electrodes. A thick layer of PMMA resist covers part of the nanotube (naturally p-type) nanotube prior to chemical doping. Image from [Avouris01].

At room temperature at which thermal noise washes out these size-induced effects, there is limited size benefit in replacing a traditional semi-conductor junction with a
molecular component because the device size is ultimately limited by the size of the lithographically defined features. If one reduces the size of the head of a hammer but not the handle size, one is not left with a smaller hammer, but rather a disproportioned one. A similar argument can be made against the expectations that replacing bulk crystalline silicon with molecular components will automatically lead to major device miniaturization. However, if each molecular component in the device is an independent and functional device, and externally/remotely addressable, the device density greatly increases.

1.1.2 – Unscaleable molecular devices in solution

Another route towards molecular machines and electronic devices is through purely supramolecular chemical syntheses of switching elements like rotaxanes and catenanes (figure 1.3) [Stoddart01] that switch their structure upon global oxidation/reduction. While these compounds are often amazing feats in chemical synthesis, their utility as functional devices is extremely limited by the fact that they are not scaleable in solution.

![Figure 1.3](image.png)

Figure 1.3 In the ([2]rotaxane 1^{+}\textsuperscript{+}), the bipyridinium-based cyclophane shuttles from one 1,4-dioxybenzene recognition site to the other depending on the pH of the solution (oxidation / reduction state of the system). Image from [Stoddart01].

Azo compounds that undergo optically driven cis-trans isomerizations are also popular molecular switches [Petty95]. Artificial oligonucleotides have been synthesized with azo-benzene compounds at the 2’ position that can be used to ratchet apart the strands in the DNA double helix by creating steric crowding upon cis-trans isomerization in the presence of UV light [Komiyama99]. Others have optically controlled phenylalanine-cyclodextrin host-guest interactions by blocking and unblocking the cyclodextrin host site with a covalently bound azo compound [Petty95]. These types of switches may lead to
interesting molecular machines, but they are volatile (with respect to memory), not bi-stable. Furthermore, the number of states that an azo switch can adopt is limited to two.
1.2 – Biological inspiration

As so aptly observed by Feynman [Feynman59], the “marvelous biological system” exhibits a staggering level of complexity that far exceeds the complexity of artificial systems at that size scale:

The biological example of writing information on a small scale has inspired me to think of something that should be possible. Biology is not simply writing information; it is doing something about it. A biological system can be exceedingly small. Many of the cells are very tiny, but they are very active; they manufacture various substances; they walk around; they wiggle; and they do all kinds of marvelous things—all on a very small scale. Also, they store information. Consider the possibility that we too can make a thing very small, which does what we want—that we can manufacture an object that maneuvers at that level!

Let us look at DNA replication as an example. DNA polymerase is an enzyme roughly 10nm in diameter that is responsible for replicating DNA from a template strand. What can this biological machine do at 1/10 the size of a single line, which is not even a functional transistor, in the current state-of-the-art Pentium processor? … It replicates at approximately 1000 base pairs per second with an error rate of one part per million [Carlson02]. This is absolutely remarkable.

It is extremely important to analyze the blueprints of biology or even consider the use of robust biological molecules as the building blocks when constructing molecular systems. Indeed, this is a trend becoming prevalent in many fields [DOE02], such as (but certainly not limited to) materials synthesis [Belcher00], computation [Winfree96], and control theory [Doyle02].

1.2.1 – Bio-inspired assemblers in the literature

Let us consider an assembler any system that can move a physical object along a linear chain. There are very few examples of assemblers in the literature, and ironically, they too are based on biological principles or are built out of biological components. The rotoxanes and catenanes mentioned previously have been called “molecular shuttles” for their ability to shuttle a molecule between two oxidation-reduction centers. These systems are largely based on the principles of metal ion coordination at biological redox centers.
There are two systems that require a 2-D substrate that are especially worth mentioning in the context of molecular assembly lines, even if they are not truly molecular by our definition. In the first system, Hopfield et al., using Photosystems I and II as an inspiration, proposed an electronic shift register memory that passes electrons down a polymeric chain by photo-induced electron transfer reactions [Hopfield89]. In figure 1.4a, a laser pulse excites an electron to the ground state of α, and then the electron decays to the ground states of β and γ, finally ending up in the ground state of the next α unit. The polymer is placed between two electrodes and bits are written by applying a large potential, thereby reducing the first α unit of the polymer. The bit is read at the drain electrode (figure 1.4b). This system is particularly important because it first introduced the notion that an object can be moved down a linear chain by pulsing a remote input. The molecular assembly line proposed in this thesis is based on this notion, as well as Hopfield’s design of a polymerized chain of repeating α-β-γ units.

In the second system, Vogel et al. [Vogel01] moved a fluorescent microtubule along a substrate of patterned kinesin motor proteins (figure 1.5). When ATP is provided, the kinesin proteins push the microtubule along the surface. The microtubule transports streptavidin-coated beads that are bound to the microtubule via binding with biotin molecules that coat the microtubule surface. The motion can also be controlled remotely by first caging the ATP molecules, and then releasing them from the cages with UV irradiation; this provides a burst in energy source and initiates the microtubule
movement. The motion stops when all the ATP is consumed (figure 1.5b). Figure 1.5c shows the displacement of the microtubule after 40 seconds; the arrows point to bound streptavidin beads as a reference for measuring the displacement. As will be discussed in section 1.2.3, this is the opposite of what happens in biology; typically, the microtubule is static, and the kinesin moves along the microtubule track.

Figure 1.5 a) A microtubule shuttle carries a streptavidin bead along a kinesin surface. b) The velocity of the movement versus time. UV pulses every 30 seconds release caged-ATP molecules and initiate the microtubule movement by providing a burst of the energy source. c) Fluorescence images of the movement after 40 seconds. The arrows point to streptavidin coated beads ("cargo") as a reference for displacement. Images from [Vogel01].

1.2.2 – Polyketide synthase and covalent hand-off

The polyketide synthase (PKS), often referred to as "Nature's assembly line," is an enzyme found in plants that produces a class of antibiotics known as polyketides. A typical polyketide synthase can synthesize very large molecules with stereoselectivity at rates on the order of $\text{pmol}_{\text{PK}} / (\text{minute} \times \mu\text{M}_{\text{PKS}})$ [Simpson98], which corresponds roughly to one perfect drug every 10 minutes per polyketide synthase from start to finish. This is an astounding rate, on the order of $10^4$ times faster than commercial drug synthesis, if one assumes it takes the pharmaceutical industry somewhere between one week and one year to synthesize molecules of similar complexity and stereoselectivity (up to roughly 10 stereoselective centers) with any reasonable yield, not including the R&D and process development time.

Figure 1.6 shows how a polyketide synthase performs its molecular assembly process. The left image delineates the assembly line model of a PKS. A linear chain is synthesized, with the addition of a two-carbon unit, known as a ketide, per reaction step. Each ketide addition occurs in a separate "module" of the PKS, and the molecule is handed-off to the next module. Each module has a minimum of three monomeric enzymes, or "domains," that are polymerized - ketosynthase (KS), chain length factor
Introduction: Biological Inspiration

(CLF), and acyl carrier protein (ACP). In the figure, 6-deoxyerythronolide (6-DEB) is synthesized linearly and then cyclized. The cyclized 6-DEB can then serve as a precursor that can be tailored into several antibiotics, in this case erythromycin A [Hopwood97].

Figure 1.6 The polyketide synthase left) Assembly-line model polyketide synthases (PKS). 6-deoxyerythronolide (6-DEB) is synthesized linearly, and then cyclized. The 6-DEB precursor can be tailored into several antibiotics, in this case erythromycin A. Image from [Hopwood97]. right) Methylation of nonribosomal peptides by methyl transferase domains in polyketide synthases, showing how the enzyme performs covalent hand-off of small molecules between domains. Image from [Walsh01].

In the right image in figure 1.6, part “a” shows the working mechanism of covalent hand-off between subunits during the methylation of nonribosomal peptides by methyl transferase domains. First, an aminoacyl thioester is catalytically methylated by the coordination of S-adenosyl methionine and the thioester by the methyl transferase (MT) and PCP domains in the n-module. The now methylated aminoacyl thioester forms an amide bond with the peptidyl chain located at the PCP domain in the (n-1)-module, and the entire peptidyl chain is handed-off to the n-module because the amide bond strength is greater than the thioester bond.

So, how can the polyketide synthase accomplish the incredible feat of assembling complex structures with extreme speed and accuracy? First, every reactive step is catalyzed. Second, it achieves stereoselectivity by coordinating the orientation of the polyketide chain within the pockets of the enzyme as it is being synthesized. Third, it performs covalent hand-off of the molecule as it is being synthesized. This essentially eliminates diffusion in the synthesis process thereby shortening the reaction time. It also contributes to the stereoselectivity of the reaction because it guarantees that the
molecules are constrained in a certain manner when they are handed-off between the pockets in the enzyme. The hand-off also serves as a feedback mechanism for the reaction completion because the molecule cannot be handed-off between sites without the reaction occurring. Therefore, in examining Nature’s assembly line, the following notion is key towards the design of a molecular assembly line:

In order to assemble something quickly and efficiently, diffusion must be eliminated by physically handing-off the object that is being assembled.

1.2.3 – Kinesin and the tethered walk

Kinesin (figure 1.7) is a motor protein that moves like a train on a filamentous microtubule track. Kinesin has several functions, like active intracellular transport of small molecules [Brady92] and stretching cells during cell division [Sharp00]. While it was originally believed that kinesin moves in only one direction like most motor proteins, from the net negative end to the positive end of the (electric) dipolar filament, it has now been shown that kinesin can move bi-directionally [Endow99].

The kinesin protein is a “two-head” protein, meaning it contains two motor regions that are bound to the microtubule track when it is not in motion. However, the “two-head” model does not preclude the possibility that only one of the heads is bound as kinesin moves along the filament; it has been shown that this is indeed the case [Vale00]. As can be seen in figure 1.7, when the protein is not in motion, both heads are bound to the filament. When the kinesin molecule moves, one head remains bound as the other head steps to the next binding site along the track. This is effectively a “tethered-walk,” and while it is not termed as such in the literature, it will be called a tethered-walk from here on out. The bias in direction (net forward motion) during the “power-stroke” comes from conformational changes induced by ATP binding at the neck between the two heads, or in other words, the head-filament interaction drives the direction of movement.

This walk is fundamentally different from two-head motor proteins like myosin in which only one is head bound at any time and the heads are independent of one another. The directional movement in myosin during the power-stroke comes from the angular rotation of its long and tilted swivel arm as ATP binds to the arm. Myosin effectively skips along
the filament. One head dissociates as the other finds another binding site, but the system is not tethered.

Figure 1.7 a) Tethered-walk in kinesin. One head is bound to the filamentous track, as the second head steps to the next binding-site on the track. b) Untethered hop in myosin in which both heads dissociate from the track as the protein moves between binding sites. Image from [Vale00].

Where does the action occur that confers directional movement in these systems? Directionality is conferred in kinesin from action that occurs between the motor heads and the filament, whereas in myosin, directionality is conferred by action in the arm and away from the filament as the entire system is dissociated. Therefore, in the context of the molecular assembly line that is being proposed (see section 1.3), kinesin movements show that:

\textit{In order to move an object along a polymer chain without dissociating the object from the polymer, a tethered-walk is required and one must control the interaction between the object and the polymer scaffold.}
1.2.4 – Further discussion on biological assembly

Polyketide synthases and kinesins were discussed in great detail because of their relevance to the design of the molecular assembly line proposed in this thesis. But, let us take a brief moment to discuss more general lessons that biological systems can teach us about the requirements for quickly assembling complex structures with extreme precision at the molecular scale.

First, the assembly should be predicated. The polyketide synthase can quickly synthesize complex antibiotics because it covalently hands-off small molecules and because each step is catalyzed. It has this ability because the assembly is predicated, which allows the structure of the enzyme to be optimized for this process. Essentially, predication is required for the optimization of the kinetics and thermodynamics of a system. Once the system is optimized, it can be evolved with precise changes. Second, feedback is required. DNA polymerase can replicate DNA with incredible speed and fidelity because it has feedback mechanisms; polymerase not only uses the template strand to direct the assembly of the growing nucleotide chain, but also checks the added nucleotide against the template strand after the addition [Glick98]. Lastly, error-correcting codes are required. Perhaps biology’s greatest example of error-correction is the Deinococcus radiodurans, an extremophilic bacterium capable of surviving incredibly extreme x-ray irradiation because it can salvage the damaged DNA strands and enzymatically reconstruct its entire genome in hours [Narumi00].
1.3 – The design of the molecular assembly line

There is no definition for a “molecular assembly line” in the literature, but in this thesis, it is defined as a system that meets the following requirements:

1) A polymer chain composed of monomers or “antennae.”
2) a. A molecular “shuttle” (e.g. an individual molecule or nanoparticle) can be moved from one arbitrary monomer position along the polymer chain to any other monomer position for the purpose of assembling some chemical moiety onto the shuttle.
   b. The shuttle can move between these positions along any arbitrary pathway determined by a programmed set of inputs. Therefore, the movement must be bi-directional and reversible.
3) The inputs into the system that move the shuttle along the assembly line must be external.
4) For the system to be truly molecular, it must be soluble in some solvent

1.3.1 – Scheme

The general scheme for a molecular assembly line is shown in figure 1.8. The molecular assembly line is comprised of a polymer with a repeating pattern of 3 concatenated monomers or antennae, -(1-2-3)_n-. The repeating pattern is shown in parenthesis in the figure. The yellow circle in the figure represents the shuttle. In the absence of any inputs, the shuttle has an equal binding affinity towards any of the antennae. Each input frequency effectively “turns off” attractive intermolecular forces between the respective antenna and the shuttle (i.e. hν₁ turns off the bond between the shuttle and antenna 1).

Assume that the shuttle is in position a, tethered between antennae 1 and 2 of the first (1-2-3) pattern (top). In order to move the shuttle from position a to position b, hν₁ is input, thereby preventing binding of the shuttle to any antenna 1. Because the shuttle is tethered to antenna 2 of this first pattern, it will be biased to toggle over to position b, between antennae 2 and 3 of the first pattern. Each time the bond between the shuttle and antenna 1 is reformed, it is broken again, but once a bond between the shuttle and antenna 3 is formed, the shuttle is in a stable position. This system is bi-directional and reversible; the shuttle can be toggled back to position a by inputting hν₁, or to position c
by inputting $h\nu_2$. The system has performed a tethered-walk. If the original position of the shuttle is known, it can be moved to any position along the molecular assembly line by any arbitrary pathway, given the correct input sequence. Therefore, the system is externally programmable.

![Diagram](image_url)

**Figure 1.8** General scheme for a molecular assembly line. The circle represents a shuttle, and each rectangle represents a different antenna. The repeating pattern is shown in parentheses.

### 1.3.2 – Applications

This system is a universal scaffold for molecular assembly processes. It moves a solid-phase support to bring reactive moieties on the support in proximity to reactive species that are located at the stable tethered positions along the backbone (sites a-e in figure 1.8). For example, the support could be a controlled-pore glass sphere with a growing DNA strand and the four nucleic acids could be located at the reactive sites; the nucleic acid of choice could be added to the growing chain by moving the bead to the desired position and then introducing polymerase into the system (figure 1.9). The molecular assembly line can also be used for building nanoelectronic devices. The simplest (1-2-3) unit shown in parenthesis in figure 1.8 is a bi-stable toggle switch.
Weck et al. have also proposed a “universal polymer backbone” (figure 1.10) that is also based on biological molecules [Weck03]. In the Weck scaffold, monomers are polymerized into co-polymers with desired periodicity by ring-opening methathesis (ROMP) reactions. Following the construction of this peptide-like backbone, an “orthogonal self-assembly” step functionalizes the scaffold into novel materials. This backbone is driven by self-assembly and designed for combinatorial materials synthesis. The molecular assembly line differs from the universal backbone proposed by Weck in that the assembly line is designed for dynamic and programmable assembly post-scaffold-synthesis.
1.3.3 – Potential shuttle/antennae interactions and backbones

There are a wide range of potential shuttle/antennae interactions that can be used to build a molecular assembly line, some of which are listed in figure 1.11. Essentially, any set of antennae with three resolvable inputs can create a molecular assembly line. As can be seen in the table, input resolvability and destructiveness, and backbone synthesis are the major issues in choosing the interaction type.

In this thesis, experimental data will be presented on the shuttle/antennae interactions in the first two proposed systems, remote electronic control of DNA hybridization state (Chapter 2), and competitive Michael addition and Norrish Type II photocleavage reactions (Chapter 4). A fully polymerized molecular assembly line has not been constructed, but routes towards them for each approach will be discussed in the respective chapters.
Introduction: Molecular assembly lines

<table>
<thead>
<tr>
<th>Interaction</th>
<th>Input</th>
<th>Pros</th>
<th>Cons</th>
</tr>
</thead>
<tbody>
<tr>
<td>Remote electronic DNA hybridization [Hamad02]</td>
<td>RF coupled into covalently attached nanocrystal antenna</td>
<td>- Highly scaleable backbone.</td>
<td>- Resolvability of input frequencies</td>
</tr>
<tr>
<td>Reversible covalent chemistry</td>
<td>Simultaneous Michael additions and wavelength selective Norrish Type II photocleavage reactions</td>
<td>- Resolvable by wavelength</td>
<td>- Destructive input (210 &lt; (\lambda) &lt; 310nm)</td>
</tr>
<tr>
<td>Conformational switching and isomerization [Petty95]</td>
<td>Blocking interactions b/w phenylalanine and cyclodextrin host by UV-induced conformational changes in azo compounds.</td>
<td>- Covalent hand-off mimics polyketide synthase</td>
<td>- Polymer backbone not easily synthesized</td>
</tr>
<tr>
<td>Electrostatic [Petty95]</td>
<td>Light driven charge transfer complexes (TCNQs).</td>
<td>- Various complexes known to absorb UV-visible light with reasonably resolvability and over the entire UV-visible spectrum.</td>
<td>- Highly basic environment is bad for assembly</td>
</tr>
<tr>
<td>Metal complex formation [Stoddart01]</td>
<td>Oxidation-reduction state of the complex forces directional movement of a metal-binding ring, such as rotaxanes and catenanes.</td>
<td>- Chemistry is well developed</td>
<td>- Charge creates highly acidic or basic local environments that are bad for assembly</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>- Global input requires that local charge be created with electrodes</td>
</tr>
</tbody>
</table>
Chapter 2: RF Biology

2.0 – Introduction to RF Biology

This chapter proposes the design of a molecular assembly line based on selective dehybridization of DNA by coupling a radio-frequency magnetic field into covalently attached nanoparticle antennae that heat the DNA, an idea that will be termed “RF Biology.” Experimental work on the monomer unit, a short DNA strand with a covalently attached superparamagnetic nanoparticle, will also be presented.

2.0.1 – Selective dehybridization of DNA

It has been shown that a DNA molecule with a 1.4nm Au nanoparticle covalently attached to it can be selectively dehybridized in a 1GHz RF field [Hamad02]. The RF energy is coupled into the nanoparticle and locally heats the biomolecule while the global temperature of the solution remains constant. Figure 2.1 shows a UV time-course spectrum (260nm) of a DNA hairpin-loop structure in which the DNA hybridization state follows that of the RF on/off cycle. The hairpin-loop is a 38mer with a seven base-pair complement. The peaks in the spectrum represent DNA in its dehybrized state; dehybridized DNA absorbs more UV light at 260nm because of a change in the dipole of the phosphate backbone, a phenomenon known as hyperchromicity.

Figure 2.1 Selective dehybridization of a DNA hairpin-loop by coupling 1GHz RF into a covalently attached 1.4nm Au55 antennae [Hamad02].
While the mechanism of local heat generation has yet to be elucidated, it is proposed that the RF inductively heats the metallic nanoparticle. Given this proposed mechanism, there is no resonance frequency at which the DNA is dehybridized, but rather, there is a threshold/basement frequency at 800MHz, above which the localized heating occurs [Hamad, UPR].

2.0.2 – Molecular assembly line based on RF Biology

As will be discussed in section 2.1, the use of superparamagnetic nanoparticles may provide the route to many channels to interface with biology because ferrofluids of superparamagnetic nanoparticles have material and size-dependent resonance absorption frequencies in the RF [Rosenzweig02].

However, let us assume that we have three nanoparticle antennae with resolvable RF absorption bands. Figure 2.2 shows an idealized molecular assembly line based on RF Biology components. Each monomer is composed of a DNA/nanoparticle assembly; the DNA strand is the same for all three monomer units (in the figure, CGTTGC with A₉ spacers). The shuttle in this case could be any non-metallic and non-magnetic solid support that is capped with DNA strands that are partially complementary to the monomer strands (in the figure, GCAACG). Because the control mechanism is localized heating, the nanoparticles must be spaced by a distance d, such that d > R, where R is the radius of a thermal radiation sphere between a point source to Tₒ. In other words, the heating must be spatially resolved between antennae. Only a (1-2-3) unit is shown for simplicity. M₁ and M₂ represent superparamagnetic nanoparticles with resolvable absorption bands.

There are many advantages in using DNA/nanoparticle assemblies as monomers for a molecular assembly line. The input is non-destructive, as opposed to optical techniques. DNA provides an excellent backbone for the molecular assembly line that can easily be scaled up to yield long assembly lines, and it is even conceivable that engineered plasmids and enzymes could synthesize the molecular assembly lines in vivo. Lastly, the ability to locally heat biological molecules and break hydrogen bonds has many potential applications in medicine and biophysics, since hydrogen bond formation is one of the main mechanisms in biological assembly and recognition.
2.1 – Introduction to superparamagnetic nanoparticles

In electronically addressing a biological molecule using nanoparticle antennae, a reasonable nanoparticle antenna must be fewer than a few nanometers in diameter in order not to dwarf or impede the biological function of the molecules, which are typically 2nm (DNA double helix) to 5nm (typical proteins) in diameter. In this size regime at room temperature, a nanoparticle composed of magnetic elements is considered superparamagnetic.

2.1.1 – Superparamagnetism

Superparamagnets are magnetic particles for which ambient thermal energy is large enough that the magnetization direction cannot be fixed by the anisotropy of the particle [Brown63]. Therefore, at ambient temperature and normal experimental time scales, a superparamagnet has no magnetization in the absence of an external field. At any given temperature, T, the probability P(t) that a particle in some metastable (magnetized) state with energy $E_1$ will be found in a state having energy $E_2 = E_1 + KV$, is given by the Arrhenius relationship in equation 2.1 – where $\tau_N$ is the relaxation time of the magnetic moment, $\nu_0$ and $\tau_0$ are pre-exponential factors (typically, $10^{-9} < \tau_0 < 10^{-11}$ s) and K is the uniaxial anisotropy energy constant that defines a free-energy barrier per unit volume.
RF Biology: Introduction

[O’Handley00]. Equation 2.1 describes a magnetization reversal mechanism of thermal activation over an energy barrier \( K_V \) between two metastable states; the transition probability between these states is a function of the ratio between the barrier \( K_V \), and thermal energy \( k_B T \). Figure 2.3 is a picture of the time-dependent magnetization, showing that the magnetization can spontaneously reverse by thermal activation over an energy barrier.

\[
P(t) = \nu_0 \exp(-K_V/k_B T) \\
\tau_N = \tau_0 \exp(K_V/k_B T)
\]  

Figure 2.3 Time dependence of magnetization direction \( \theta \) with respect to an applied field, \( H_a \). \( N(t) \) represents the number of particles in a particular energy state with respect to time. The magnetizations can spontaneously reverse by thermal activation over an energy barrier. Image from http://www.essrl.wustl.edu.

Below the “blocking temperature” at which the thermal energy of the system is insufficient to demagnetize the nanoparticle by spontaneous reversal of magnetization, thermal fluctuation is effectively “blocked” and the magnetization remains in a metastable state. Above the blocking temperature and at normal experimental times, a superparamagnet shows no hysteresis in its M-H curve.

Superparamagnets interact with magnetic fields differently at various frequencies. In a static (DC) field, if the particle is free to rotate, it acts as a magnetic dipole and the nanoparticle aligns with the field because it can reduce its energy by aligning its magnetization with the easy (anisotropy) axis and the applied field. In a sufficiently concentrated solution (ferrofluid) or viscous medium, the nanoparticles will form chains because of dipole attractions between particles. In AC fields, the heating mechanisms are due to Brownian losses (kHz), Neél losses (MHz), or ferromagnetic resonance (GHz). These losses are described below. Although, for reasons that will be discussed
in this section, heating by Neél losses was the heating mechanism of choice, it is important to understand each mechanism for future work in designing systems that interface biological molecules with nanoparticle antennae.

### 2.1.2 – Low-frequency RF losses: Brownian relaxation

At $\nu \ll 1/\tau_N$, Brownian rotation of the entire particle is the predominant loss mechanism. The entire particle acts as a dipole that tracks the field with the relaxation time, $\tau_B$. The torque ($T$) on the particle and the power dissipation ($P$) by viscous drag models [Thompson98] are given in equations 2.3 and 2.4:

\[
\tau_B = 3V\eta/k_BT \tag{2.2}
\]

\[
T = \mu_o \pi R^3 H_o^2 \tag{2.3}
\]

\[
P = 2N_{\text{particles}} \mu_o \pi^2 R^3 H_o^2 \nu \tag{2.4}
\]

where, $\eta = $ solvent viscosity, $R = $ particle radius, $H_o = $ applied field strength, and $\nu = $ frequency of the AC field.

Power dissipation by viscous drag is the primary cause of heating in magnetic fluid hyperthermia (MFH), a method of destroying tumor cells in low-frequency RF fields by loading them with superparamagnetic nanoparticles. However, the cellular environment is extremely viscous compared to water, in which a molecular assembly line using a DNA backbone would have to be. Furthermore, as can be seen in equation 2.4, it is the size of nanoparticle that primarily affects the power dissipation, and therefore it would be difficult to create localized heating in one antenna (by torque) without influencing all other magnetic nanoparticles in this frequency regime.

### 2.1.3 – High-frequency RF losses: Neél relaxation

At frequencies on the order of $1/\tau_N$, Brownian motion is essentially frozen out as a mechanism for aligning the particle magnetization with the field. Therefore, the magnetization rotates within the particle to track the field. Losses by this mechanism are known as Neél losses. In a ferrofluid, the magnetization relaxation time $\tau_{\text{eff}}$ depends on
RF Biology: Introduction

$\tau_B$ and $\tau_N$, but as can be seen in equation 2.5 and figure 2.4a, $\tau$ is essentially determined by $\tau_N$ for small particles.

$$1/\tau_{\text{eff}} = 1/\tau_B + 1/\tau_N \quad (2.5)$$

$$\chi(\omega) = \chi'(\omega) - i\chi''(\omega) \quad (2.6)$$

$$\chi(\omega) = \chi_o/(1+\omega^2\tau_N^2) - i\omega\tau_{\text{eff}}\chi_o(1+\omega^2\tau_N^2)$$

where $\chi_o = \text{initial susceptibility}$

$$P = \pi \mu \chi_o H_o^2 \nu (2\pi \nu \tau) / [1 + (2\pi \nu \tau)^2] \quad (2.7)$$

$$\Delta T = P\Delta t/c \quad (2.8)$$

where $c = \text{specific heat}$

Heat can be generated by Neél losses by the phonons created in the magnetization rotation process. As $\omega \tau$ approaches unity, where $\omega$ is the angular frequency, $2\pi \nu$, the imaginary susceptibility ($\chi''$) reaches a local maximum and the real susceptibility ($\chi'$) passes through a local inflection point (equation 2.6, figure 2.4b) [Fannin02]. The power dissipation also approaches a maximum as $\omega \tau$ approaches unity; the equations for power dissipation and temperature change are given by equations 2.7 and 2.8.

Figure 2.4 a) Ferrofluid relaxation time $\tau_{\text{eff}}$ as a function of $\tau_B$, $\tau_N$, and diameter. Image from [Rosensweig02]. b) Complex susceptibility curves of 8.4nm cobalt nanoparticles. The peak at 25MHz corresponds to maximum power dissipation from Neél losses. Higher frequency resonances are result from ferromagnetic resonance. Image from [Fannin02].
2.1.4 – Microwave losses: Ferromagnetic resonance

At microwave frequencies (GHz), the magnetization cannot track the field by Neél losses. Instead, the magnetization and physical orientation of the particle are pinned along the anisotropy axis by the longitudinal component of the AC field. As the drive frequency approaches the precession frequency of the magnetic moment about the anisotropy axis, $\omega_o$ [Fannin99], the magnetic moment can be tipped by the perpendicular component of the field, a phenomena known as ferromagnetic resonance. In FMR, the angular resonant frequency is:

$$\omega_o = \gamma(H_A + H)$$

$$H_A = 2K/M_s$$

where, $\gamma$ = gyromagnetic ratio, $H_A$ = internal/zero field of a particle, $H$ = applied static field, $K$ = uniaxial anisotropy constant, and $M_s$ = saturation magnetization.

An analogy can be made between FMR and nuclear magnetic resonance (NMR), in the sense that in FMR, the crystalline anisotropy axis acts as an internal static field ($H_A$) about which the magnetic moment precesses, as opposed to the external static field in NMR. This can be clearly seen in equation 2.9; there exists a natural precession frequency in the absence of an external static magnetic field, and the frequency increases linearly with applied static field, $H$. FMR in the absence of an applied static field is known as “zero-field resonance.”

2.1.5 – Discussion on heating mechanisms

In determining the heating mechanism of choice, several factors must be considered, and each mechanism of heating has its own pros and cons.

First, how do AC fields affect biological molecules in the absence of a nanoparticle? High-frequency fields may cause tissue damage or induce “anomalous backbone vibrations” in DNA [Nuss97]. Second, how do AC fields affect water in the absence of a nanoparticle? The issue of heating water with microwaves is troublesome in the context of the molecular assembly line and more general biomedical and biophysical contexts because it globally affects every molecule in solution. Third, how tight are the resonance
absorption bands of various nanoparticles in the associated frequency range? Developing a set of nanoparticle antennae is necessary for building a molecular assembly line based on RF Biology. It is also desirable for biophysical studies because it would allow one to control many biological molecules independently of one another. Lastly, is the physical process by which heat is generated conducive towards building a molecular assembly line?

Low-frequency Brownian losses are attractive because the AC field will not damage biomolecules or microwave water. However, the ability to find resolvable operating frequencies is an issue. The fact that the entire particle rotates is not desirable for building a molecular assembly line because the rotation of a nanoparticle antenna would affect the entire system.

Utilizing Neél losses is an attractive option because the effects of the AC field on biological molecules and water are limited in the MHz frequency regime. The nanoparticle RF absorption bands are reasonably resolvable. The heat generation is fairly independent of nanoparticle orientation.

Heating by ferromagnetic resonance is an interesting option, primarily because FMR bands are very sharp. However, one runs the great risk of damaging biological molecules and microwaving water at such frequencies.

Given the task at hand, the experimental work in this thesis was directed towards coupling RF in the MHz frequency range, or heating superparamagnetic nanoparticles by Neél losses.
2.2 – Experimental

All reagents were of the highest purity available and were used as received from Sigma-Aldrich unless otherwise indicated. Reactions under nitrogen were performed in a MBraun UniLab glove box unless otherwise noted.

High-resolution TEM micrographs were attained using a JEOL 2010 TEM on copper/Formvar grids from Ladd Research Incorporated. FTIR data was taken using a Digi-Lab Excalibur FTS 3000 (KBr, 64 scans, 4 cm\(^{-1}\) resolution). Fluorescence spectra were obtained using a Jovin-Yvon Spex Fluoromax-3 spectrophotometer (3mm x 3mm quartz cuvette, 1nm slits, 0.5s integration).

2.2.1 – Cobalt nanoparticle synthesis

Synthesis of cobalt nanoparticles was performed under nitrogen. Cobalt chloride (CoCl\(_2\), 1.0mmol) and 11-aminoundecanoic acid (2.0mmol) were added in octyl ether (20mL, 0.05M\(_{Co}\)) and heated to 100°C. Tri(octyl)phosphine (3.0mmol) was added, and the solution was heated to 160°C. 1.0M Super-Hydride in THF (2.0mL, 2.0mmol) was added and the solution was stirred vigorously for 20 minutes; the solution color immediately changes from “cobalt blue” to black. Additional Super-Hydride was added (1.0mL, 1.0mmol), and the solution was stirred for 20 more minutes. The reaction was then slowly cooled to room temperature and then removed from the nitrogen atmosphere. The nanoparticles were purified by centrifugation in pure ethanol (approximately 3mg/mL) for 2 hours at 6000g followed by decantation, two times, and then air-dried.

2.2.2. – DNA/nanoparticle assemblies (2)

The DNA hairpin-loop (1) (“stem-loop” or “molecular beacon”) was synthesized by Genset Oligos. The terminal nucleotides made up a 7-base-pair stem, with a 24-base-pair loop. The fluorescein dye, FAM, was appended at the 5’ end, and the quencher, Dabcyl, at the 3’ end. The eighteenth base, denoted as T’, was a thymine with an internally modified free amine. The sequence of the oligonucleotide is (5’ → 3’) given below with the portion in parenthesis representing the “loop” of the hairpin-loop structure:

FAM – GCGCCCT – (AAACTGGTGT’GGAAATGCGTCATG) – AGGGCGC – Dab
Oligonucleotide dehybridization causes an increase in fluorescence, as the efficiency of Dabcyl quenching of FAM fluorescence is distance dependent. The idealized structure of the hairpin-loop and the scenario of increased fluorescence upon dehybridization are shown below in figure 2.5a.

**Figure 2.5 a)** Idealized structure (left) and increased fluorescence upon melting (right) of the DNA hairpin-loop. Image modified from [Bonnet99]. **b)** Coupling and purification strategy of covalent attachment of cobalt to DNA hairpin-loop.

Upon receiving the oligonucleotides, a 30µm stock solution was made in DMSO. To obtain carboxylic acid functionalized hairpin-loops, 100µL of the stock solution (3nmol) was shaken overnight with 100µL chloroacetic acid in DMSO (300µM) (10x, 30nmol) at room temperature, using standard nitrogen techniques. The oligonucleotide was purified by overnight ethanol precipitation and then redispersed in 500µL DMSO. A tenfold excess of cobalt nanoparticle (3mg, approximately 30nmol) and a large excess of ethyldiethylaminopropylcarbodiimide (EDC) were added and the solution was shaken overnight. The sample was lyophilized, and then purified by overnight ethanol precipitation.
precipitation. The DNA/nanoparticle assembly (2) was recast in 1x PBS buffer to make a 1µM solution. The coupling scheme is shown in figure 2.5b.

2.2.3 – Application of RF to DNA/nanoparticles

Typical fluorescence spectra were averaged four times with, $\lambda_{\text{exc}} = 495$nm.

In order to apply RFMF in the fluoremeter with good global temperature control, a customized cell holder (figure 2.6) had to be constructed to replace the standard cell holder in the spectrophotometer. The cell holder was designed using Rhinoceros 3D modeling software and printed with ABS plastic using a Stratsys 3-D printer. The solenoid (Cu wire, 7 turns, d = 3cm, h = 3cm) was potted using epoxy and painted black to eliminate fluorescence from the polymer. A 0.5cm gap was placed between the solenoid and the cuvette. Water-cooling was provided between the solenoid and sample on the two sides that were not in the light path of fluoremeter to limit radiative heat transfer from the solenoid; the cooled water (15°C) was pumped through Master-Flex tygon tubing with a peristaltic pump, and the top portion of the cuvette that was not in the solenoid core was wrapped with tubing. The entire apparatus was cooled with a small cooling fan (Nidec Gamma 28).

RF was applied with a HP8648C signal generator coupled with a linear amplifier (+35dBm, 1mW input saturation, 4W output saturation). Because the coil was not impedance matched to 50Ω over all frequencies, the signal generator was driven at maximum output (+14.5dBm) in order to ensure a 4W signal from the amplifier. Prior to RF biology experiments, the power from the generator was increased from 0.0-14.0dBm,
and power after the amplifier was monitored by use of a bi-directional coupler (e-meca) and spectrum analyzer (Agilent E4407B); if the power output from the amplifier saturated before 14.5dBm, a 4W output was assumed. The calculated field strength was 8.2mT.

2.2.4 – Enhanced microwave heating of non-polar solvents

2mL of commercially available ferrofluid of 10nm Fe₃O₄ nanoparticles in a hydrocarbon carrier (FerroTec USA) was dissolved in 8mL hexanes, yielding a 1% volume fraction ferrofluid. The solution was microwaved for two minutes in a commercially available microwave oven (Sharp R214-FW) with a center frequency of 2.45GHz.
2.3 – Results and Discussion

2.3.1 – Cobalt nanoparticle synthesis

The $\varepsilon$-cobalt nanoparticle (NP) synthesis was based on methods reported by Murray [Murray99]. $\varepsilon$-Co or epsilon-phase cobalt is a soft magnetic material whose structure resembles that of the $\beta$-phase of manganese. It has a reduced anisotropy from that of HCP cobalt, but can be converted into HCP cobalt by annealing at high temperatures. The low magnetic dipole interactions between particles in this phase during synthesis allows for slower and more controlled nanoparticle growth rates of small superparamagnetic nanoparticles [Murray99][Alivisatos01].

The phosphine provides steric hindrance to slow nanoparticle growth during Super-Hydride reduction. Phosphines weakly bind to the cobalt ions, but are incapable of reducing oleic acid-chelated cobalt ions to form nanoparticles. In general, longer alkyl chains on the phosphine lead to smaller nanoparticles. Murray reports the use of tri(butyl)phosphine yields 7-11nm nanoparticles, and tri(octyl)phosphine yields 2-6nm nanoparticles.

An excess of ligand was used, and the reaction was run at a lower temperature to obtain small nanoparticles. Figure 2.7 shows TEM micrographs and the size distribution of the nanoparticles as synthesized; nanoparticles of 1-5nm were synthesized, with a mean particle size of 2.8nm ($\sigma = 0.7$nm). Because of the induced dipole forces between superparamagnetic nanoparticles during synthesis, it is extremely difficult to achieve one uniform size, especially since the induced dipoles are likely very large when stirring the solution with a magnetic stir bar. Despite reasonably good coarse size control during synthesis, size-selective centrifugation or flocculation is required for fine size selectivity and should be employed in the future. Future work should also employ a mechanical stirrer to lessen induced dipolar attractions during the ripening/growth process, thereby yielding a tighter size distribution in the initial synthesis step.

For the nanoparticles to be covalently bound to biological molecules, terminal reactive functional groups are required, and the nanoparticle must be soluble in aqueous and polar organic solvents. Synthesis with dioic acids (HOOC-(C$_n$)-COOH) led to very large,
gray precipitates, presumably because of bridging between particles, even in the presence of an excess of dioic acid. The aggregates were insoluble in all solvents, even with 15 minutes sonication. TEM showed particle sizes of 100-500nm.

\[ \text{Figure 2.7 a) Size distribution of cobalt nanoparticles obtained from TEM micrographs, with } d = 2.8 \pm 0.7 \text{nm. b) TEM micrograph of cobalt nanoparticles. Scale bar } = 10 \text{nm. c) TEM micrograph of 4nm single crystalline cobalt nanoparticle.} \]

Therefore, 11-aminoundecanoic acid was the preferred ligand, and the coupling reaction of choice between the biomolecule and nanoparticle was for the biological molecule to have carboxylic acids and nanoparticles to have terminal amines. Direct synthesis with 11-aminoundecanoic acid was preferred over the more common strategy of synthesis with long alkyl chain carboxylic acids, like oleic acid, followed by ligand exchange, in order to limit the number of purification steps required and to ensure the percent composition of the terminal group since ligand exchange is a stochastic process.

The nanoparticles were soluble in almost all polar solvents, except for acetonitrile. However, they easily formed aggregates because of the magnetic dipole and hydrogen-bonding interactions between them. The solubility was greatly increased by the addition of a strong acid like 1M HCl.
Figure 2.8 Spectroscopic analysis of cobalt nanoparticles. a) FTIR spectrum of cobalt nanoparticles. b) UV-visible spectrum. The dip around 650nm is an artifact of the spectrometer lamp.

Figure 2.8a show the FTIR spectra of the nanoparticles as synthesized. The nanoparticles show a very broad carbonyl region with peaks at 1581 cm\(^{-1}\) and 1425 cm\(^{-1}\) (carboxylate stretch), and 1500 cm\(^{-1}\) (C-H bend). The primary amine absorption is most likely buried in the region. Importantly, there is no peak above 1581 cm\(^{-1}\), meaning that all the ligands are bound to the nanoparticle by carboxylate bonds and the nanoparticle is amine-capped. Also, the absence of the carboxylic acid peak is fairly good evidence
for the removal of excess ligands. The UV-Visible spectrum in figure 2.8b shows no surface plasmon peaks, which is characteristic of very small nanoparticles. The dip in the spectrum around 650nm is an artifact of the spectrometer lamp.

Despite there being two functional groups on the ligand that are capable of binding to cobalt ions, acids and amines, only the amines are terminal. The ligand is a zwitterion, meaning that at neutral pH, the amine deprotonates the acid, leaving a positively charged ammonium and negatively charged carboxylate; since the metal is cationic, the anionic carboxylate, not the ammonium, binds to the cobalt core. This has been shown in the literature during cobalt nanoparticle synthesis in the presence of oleyl amine and oleic acid [Dumestre02]. It is important that the phosphine and Super-Hydride are added after the ligand is already bound to the metal. If the reducing agents were added with the ligand, the ligand would not be zwitterionic, and this might perhaps make it possible for the amines to bind to the cobalt, leaving mixed terminal functionalities.

2.3.2 – DNA/nanoparticle assemblies

Acrylamide gel electrophoresis of the DNA/nanoparticle assemblies and unlabelled DNA hairpins would indicate that the labeling efficiency was nearly quantitative (figure 2.9). No DNA/nanoparticles can be seen after staining; since the DNA/nanoparticle is net positively charged, it does not migrate towards the positive electrode of the gel, but rather it migrates out of the gel towards the negative electrode. Monofunctionalization (1 nanoparticle : 1 hairpin linkage) was only assumed given the large excess of nanoparticles during the coupling step. Current methods of synthesizing monofunctional nanoparticles based on surface chemistry instead of stoichiometry [Sung, UPR] should be employed in future work.

As figure 2.9b shows, the melting curve of the DNA/nanoparticle assembly is different from the unlabelled DNA hairpin-loop. The melting point is lowered by induced mechanical stress or steric hindrance from the nanoparticle. Also, the fluorescence is greatly quenched by coupling with nanoparticle surface plasmons [Dubertret01]. To account for the effects of destabilization and fluorescence quenching by covalently attached nanoparticles, effective temperatures obtained from fluorescence data are calculated from the melting curves of each individual sample. Therefore, the samples are self-consistent. From here on, the calculated temperature based on FAM emission
will be called “effective temperature.” Future work should use DNA hairpin-loops with
dyes that emit light at much higher wavelengths to limit quenching.

(1,7) = DNA stack, (2) DNA hairpin-loop with no nanoparticle, (3) Cobalt nanoparticle, (4) DNA hairpin-loop +
non-covalent cobalt nanoparticle, (5) DNA hairpin-loop with covalently attached cobalt nanoparticle
precipitate from ethanol precipitation, and (6) DNA hairpin-loop with covalently attached cobalt nanoparticle
supernatant from ethanol precipitation. The net positive charge of DNA with covalently attached
nanoparticle causes the assembly to leave the gel. The absence of a spot after staining indicates covalent
linkage. The nanoparticle does not pull DNA into the ethanol layer, as evident in lane 6.

Figure 2.9 a) Gel electrophoresis of DNA/nanoparticle assemblies (2), unlabelled DNA hairpin-loop (1), and
a mixture of (1) and cobalt nanoparticles. b) Melting curves of DNA with no cobalt nanoparticle (1) (black
circle), non-covalent cobalt nanoparticle (green triangle), and covalently attached cobalt nanoparticle (2) (red
square).
2.3.3 – Application of RF to DNA/nanoparticles

As can be seen in figure 2.10, the global temperature of 1x PBS buffered water does not change in the RF apparatus with temperature control over one hour continuous irradiation. In figure 2.10a, the frequency was swept from zero to 1 GHz, in 50MHz steps every three minutes. From here on, global heating of the solution from the heat generated by the coil or RF absorption by 1x PBS buffer will be termed “coil heating.” Continuous irradiation of 1x PBS buffered water over one hour at 300MHz also showed no evidence of coil heating (figure 2.10b). Global heat created by the solenoid was greatly limited to a temperature variation of +/- 4°C, as measured by a thermocouple (Omega). Even though water-cooling was provided between the cuvette holder and the solenoid, it was extremely important that the top portion of the cuvette was wrapped with tubing and the entire apparatus was air-cooled because the greatest source of coil heating was the heating of quartz cuvette.

![Graph](image)

Figure 2.10 a) Global temperature of 1x PBS buffered water under constant RF irradiation over 1 hour. The frequency was increased by 50MHz every 3 minutes from 0-1GHz b) Global temperature of 1x PBS buffered water under 300MHz irradiation for one hour.

Figure 2.11 shows the fluorescence spectra of FAM emission (4 scans averaged over 200 seconds) in a 300MHz field. The increased intensity of FAM emission is evidence of selective dehybridization of (2) at 300MHz versus (1), which has no nanoparticle antenna that can couple RF. There is a dip in the spectrum of (2) around 505nm and a hump around 512nm that are not typically present in the FAM emission spectrum. These aberrations are likely due to surface plasmon (figure 2.9b) coupling between the
metal nanoparticle and dye emission [Dubertret01]. The effective temperature change due to RF is $\Delta T = +18^\circ C$.

Figure 2.11 Selective hybridization of DNA hairpin-loop with covalently attached 2.8nm cobalt nanoparticle antenna. a) Fluorescence spectra of (2) in a 300MHz field for 200 seconds (red). b) (1) in 300MHz field under the same conditions as in (a).

Figure 2.12a shows the frequency response of the DNA/nanoparticle assemblies (2) from 100MHz – 1GHz, measured in 25MHz steps; y-axis data points in figure 2.12a are the fluorescence emission values at 517nm taken from spectra obtained under the same...
RF Biology: Results and Discussion

conditions as in figure 2.11. The frequency dependent selective dehybridization of the DNA hairpin-loop was not related to the field strength at that frequency, as can be seen by the reflected power spectrum of the coil from 100MHz to 1GHz, as measured by an Agilent 8417E RF network analyzer (figure 2.12b). Even though slight RF induced dehybridization occurred at all frequencies measured, the response was fairly resonant, with a Gaussian fit showing a full-width-half-maximum (FWHM) of approximately 40MHz.

\[
\text{FWHM} = 40 \text{MHz}
\]

Figure 2.12 a) Maximum FAM emission vs. frequency from (2). The Gaussian fit shows that RF coupling into the nanoparticle antennae is resonant, centered at 306MHz with FWHM = 40MHZ (or \( \sigma = 20\text{MHz} \)). b) Reflected power spectrum of the coil; the response of (2) was not related to the field strength of the coil.

While the response was not truly “resonant” like quantized UV electron transitions, a 40MHz FWHM would certainly be sufficient for building a molecular assembly line based
on RF Biology. The result would lead one to believe that another nanoparticle antenna of different composition could selectively dehybridize an oligonucleotide at another resolvable frequency below 800MHz (the cutoff of Au$_{55}$), meaning that the RF Biology approach is a reasonable one towards constructing a molecular assembly line.

In the more general biomedical and biophysics contexts, it is extremely convenient that (2) responds well below microwave frequencies. 300MHz is a reasonable operating frequency for biophysical studies and biomedical applications considering that current NMR spectrometers / MRI machines usually operate from 200 – 500MHz. It is desirable in biomedical applications that the lowest frequency possible be used in order to limit undesirable tissue damage [Hergt98] and to prevent inducing random backbone vibrations in oligonucleotides and proteins [Nuss97].

This work assumed a steady-state heating model. The effective temperature was calculated from measurements that had to be made by averaging many scans over the time span of minutes primarily because of instrument noise and the acquisition time of the spectrometer. No matter what the application, but especially if one wishes to build a molecular assembly line based on RF Biology, RF would ideally be applied for as short a time as possible and not for minutes as in this study. Future work should include single molecule kinetic studies with known heat transfer constants; to test the hypothesis that the system is one-dimensional transient conduction, the conduction and convection rates must be known for interfaces between DNA/nanoparticle/solution as well as the interface between the solution/cuvette/cooling mechanisms. The spectrometer should be able to acquire data at a frequency greater than the applied RF frequency to observe the time evolution of heat generation and selective dehybridization.

While global temperature changes were monitored with a thermocouple before and after the RF was applied, there were no direct temperature measurements during RF application because electronic thermometers (thermocouple and IR thermometer) could not function properly in the presence of the RF. The physical constraints of the spectrometer made it impossible to use a non-electronic thermometer. Therefore, future work should also employ an internal temperature standard, such as another DNA hairpin-loop with a different fluorophore with no covalently attached nanoparticle antennae.
2.3.4 – Enhanced microwave heating of non-polar solvents

Figure 2.13 shows the global temperature of a 1% volume fraction ferrofluid of 10nm \( \text{Fe}_3\text{O}_4 \) nanoparticles in hexanes and pure hexanes heated in a commercial microwave oven. The global temperature of the ferrofluid increases at a much faster rate than pure hexanes as a result of coupling energy into the nanoparticles by FMR. The global temperature increase in the pure hexanes sample, which is non-polar, is likely a result of microwave heating of the vial. Fannin et al. have shown that 10nm \( \text{Fe}_3\text{O}_4 \) nanoparticles in a hydrocarbon carrier (1.8% volume fraction in Isopar M) exhibit a zero-field FMR peak around 2.2GHz [Fannin01]. Even though a commercial microwave oven is designed for a center frequency of 2.45GHz, its spectrum is typically very broad and includes electromagnetic waves at 2.2GHz.

![Figure 2.13](image)

*Figure 2.13 Enhanced heating of hexanes solution by ferromagnetic resonance heating of 10nm \( \text{Fe}_3\text{O}_4 \) nanoparticles.*

FMR provides an interesting route towards localized heating of biomolecules through nanoparticle antennae because the resonance frequency bands are typically very sharp, and thus may provide many resolvable operating frequencies. However, as mentioned in section 2.1.5, microwave frequencies may cause damage to biological molecules, often by anomalous backbone vibrations [Nuss97]. Since the zero-field FMR frequency is primarily determined by nanoparticle size and composition, the resonant frequency for nanoparticles of desirable size to interface with biology would likely be far greater than...
2.2GHz, which is undesirable due to the efficient microwave absorption by water at those frequencies.
3.0 – Introduction to DC Biology

This chapter discusses the control of secondary protein structures in the presence static magnetic fields. The novel synthesis of magnetic wires that are exactly one molecule wide is also discussed.

3.0.1 – Control of biological molecules using static magnetic fields

In recent years, much work has been conducted on the manipulation of biological molecules and cells in the presence of static magnetic fields using superparamagnetic nanoparticles. Magnetic tweezers capable of applying and measuring piconewton forces are often used for biophysical studies like determining the relaxation mechanism of DNA supercoils by topoisomerase, or biological sensing studies like real-time PCR monitoring [Croquette00] (figure 3.1). The combination of magnetic fields and fluid flow has been used to separate cell/magnetic bead conjugates from unlabelled cells [Bacri00].

In the presence of a magnetic field, the induced dipolar attractions (N-S) and repulsions (N-N, S-S) cause superparamagnets to form chains along their anisotropy axes in a sufficiently viscous environment in which diffusion is limited. Chain formation and crystalline alignment will lead to observable room temperature hysteresis in a M-H curve, as can be seen in figure 3.2, which show M-H curves of the single-molecule magnet (section 3.0.2) Mn12-acetate (4), with and without field-alignment.
Simple chain formation of superparamagnets in the presence of a magnetic field can potentially be used to alter secondary protein conformations, as shown in figure 3.3. In the figure, two superparamagnetic nanoparticles that are covalently linked to the ends of an \( \alpha \)-helix are separated by the helix by a distance \( r_1 \) in the absence of a magnetic field. In the presence of a magnetic field, if the magnetic dipole attraction is large enough, the nanoparticles will exert enough force (torque) to induce an \( \alpha \)-helix/random-coil transition in the protein secondary structure, such that the nanoparticles will be separated by a shorter distance \( r_2 \). These changes would be observable by circular dichroism (CD) or SQUID magnetometry measurements. The figure also shows how an organic linker molecule of length greater than or equal to \( r_2 \) can be used to covalently bridge the nanoparticles once the induced conformational change has occurred, allowing one to lock the protein in its less thermodynamically favored state in the absence of a magnetic field and effectively, “turn off” the protein permanently.

Because the input is DC, there is only one possible input, making this control of protein secondary structures useless towards building molecular assembly lines. However, it was explored nonetheless since the ability to control protein secondary structures would be an extremely useful tool in biophysics.
3.0.2 – Single-molecule magnets: Special superparamagnets

A special class of superparamagnets are dodecanuclear manganese species with the formula \([\text{Mn}_{12} \text{O}_{12}(\text{RCO}_2)_6(\text{H}_2\text{O})_4]\bullet(\text{RCO}_2)_2(\text{H}_2\text{O})_4\), often called single-molecule magnets or SMMs [Lis80] [Sessoli93]. Figure 3.4 shows the structure of a typical SMM. It is comprised of 8 Mn(III) ions that form a crown around 4 Mn(IV) ions that make up an inner cube. The MnO core is less than 1nm in diameter, and the total hydrodynamic radius is less than 2nm.

![a) Typical Mn₁₂O₁₂ core of single-molecule magnets with the formula Mn₁₂O₁₂(RCO₂)₁₆. The solid black lines represent Mn-O bonds that lie on the Jahn-Teller axis. These ligands can be selectively exchanged due to the decreased bond energy that results from their distorted shape. b) Structure of Mn₁₂O₁₂(RCO₂)₁₂(NO₃)₄ showing the sites of selective ligand abstraction, represented by the solid black lines. Images from [Christou01].](image)

SMMs have large anisotropy energies, almost that of bulk materials, due to Jahn-Teller distortions in the coordination of the carboxylate ligands about the Mn ions. Their long Néel relaxation times \((\tau_o \sim 10^{-7}\text{s})\) result from geometric spin frustration between the crown Mn(III) and cube Mn(IV) ions.
These species are very attractive nanoparticles to covalently attach to biological molecules. First, they are molecular (as opposed to colloidal) and thus, have an exact and small size. Second, it has been shown that the four carboxylate ligands with both oxygen atoms lying on Jahn-Teller axes can be selectively abstracted [Christou01] (figure 3.4b) because of the decreased metal-ligand bond strength that results from the Jahn-Teller shape distortion; therefore, it is possible to guarantee covalent attachment at a particular site and along a particular crystalline axis.

The initial interest in these species was in their potential use as antennae for selectively dehybridizing DNA with RF, but they could not generate enough heat to do so. However, they may be suitable for controlling peptide conformations, as described in section 3.0.1. Furthermore, they make interesting precursors in the synthesis of molecular magnetic wires that are exactly one-molecule wide, or rather a pearl necklace of covalently linked superparamagnets that are aligned along one axis by purely chemical means.
3.1 – Experimental

All reagents were of the highest purity available and were used as received from Sigma-Aldrich unless otherwise indicated. Reactions under nitrogen were performed using standard techniques. High-resolution TEM micrographs were attained using a JEOL 2010 TEM on 200 mesh copper/Formvar grids from Ladd Research Incorporated. FTIR data was taken using a Digi-Lab Excalibur FTS 3000 (KBr, 64 scans, 4cm⁻¹ resolution). Room temperature M-H curves were obtained using a Lakeshore Scientific vibrating sample magnetometer (VSM) in epoxy. Circular Dichroism spectra were obtained with an Aviv Instruments Circular Dichroism Spectrophotometer Model 202 (3s collection time, 1nm per 1s steps). EDAX analysis was performed on an FEI DB-235 focused-ion-beam microscope (FIB) equipped with an EDAX Phoenix analyzer.

3.1.1 – Peptide/nanoparticle assemblies

Peptides were synthesized at the MIT Biopolymers Laboratory using standard FMOC/DDE synthetic techniques on chlorotrityl resins, and were received cleaved from the resins. The sequence was (AAKK)₈A (1). Upon receiving the peptides, 0.05g/mL (~0.02mmol/mL) stock solutions were made in DMF. Cobalt nanoparticles were synthesized as described in section 2.2.1. Mn12-tuBu/nitrate was synthesized as described in section 3.1.3.

First, 25µL piperidine was added to 100µL stock solution (2µmol peptide, 20% piperidine v:v) and shaken for 15 minutes at room temperature, and purified by cold ether precipitation (90% v:v, -20°C). The N-terminus was converted to a carboxylic acid by treating the peptide with 100µL of 1M chloroacetic acid in DMF (0.1mmol, 500x), shaking the solution for 30 minutes, and purifying by ether precipitation. The peptide was then recast in DMF.

18mg cobalt nanoparticles (~10x) and 38mg EDC (100x) were added to 1µL peptide (20nmol). 99µL DMF was then added, and the solution was shaken overnight, followed by purification by ether precipitation. DDE side chain protecting groups were removed by treatment with 100µL of 2% (v:v) hydrazine monohydrate in DMF for 30 minutes. After purification by ether precipitation, the resultant cobalt labeled peptide (2) was recast in 1x PBS.
Mn12-tBu/nitrate labeled peptides (3) were prepared as described above, except 5mg (6) (20x) was added with no coupling agent, in place of cobalt nanoparticles and EDC.

3.1.2 – Field-induced changes in secondary protein structures
Circular dichroism measurements were made in the presence of a magnetic field by placing a small permanent magnet under the cuvette; the position of the cuvette in the spectrometer was not altered by the presence of the magnet.

3.1.3 – Synthesis of molecular magnets and magnetic wires
All single-molecule magnet syntheses were verified by comparison of FTIR spectra with reported literature spectra.

*Mn12-acetate (4)* [Mn$_{12}$O$_{12}$(CH$_3$CO)$_2$$_{16}$(H$_2$O)$_4$]
Synthesis of (4) was based on literature methods [Lis80]. 4.0g (16.3mmol) [Mn(CH$_3$CO)$_2$•4H$_2$O] was dissolved in 40mL 60% (v/v) acetic acid (aq). 1.0g KMnO$_4$ (6.4mmol) (Mallinckrodt) was added and the solution was stirred for two days at room temperature. The red powder was collected by vacuum filtration and washed with copious amounts of acetone and THF. The disordered acetate ligands in [Mn$_{12}$O$_{12}$(CH$_3$CO)$_2$$_{16}$(H$_2$O)$_4$]•(CH$_3$O)$_2$ were removed by toluene azeotrope (2 x 75mL) followed by vacuum filtration and washing with acetone, and then the process was repeated.

*Mn12-tBu (5)* [Mn$_{12}$O$_{12}$(tBu-CH$_2$CO)$_2$$_{16}$(H$_2$O)$_4$]
Synthesis of (5) was based on literature methods [Christou01]. 2.0g (1mmol) of (1) was dissolved in 50mL toluene and treated with 2mL tert-butylacetic acid. Toluene and acetic acid (azeotrope) were removed under vacuum. 50mL of toluene was added and removed by evaporation. After the entire process was repeated, the slurry was dissolved in 50mL toluene and filtered. The collected solid (5) was washed with hexanes and dried under vacuum.
Mn12-tBu/nitrate (6) [Mn_{12}O_{12}(tBu-CH_2CO_2)_12(NO_3)_4(H_2O)_4]
Synthesis of (6) was based on literature methods [Christou01]. 0.1g (0.04mmol) of (5) was dissolved in 5mL acetonitrile and treated with 900µL of 0.226M HNO_3 (in acetonitrile). The slurry was stirred for one hour. 2.5mL nitromethane was added, and the solution was filtered. The collected solid was washed with ether and dried under vacuum.

Molecular wires based on Mn12-acetate (7)
0.05g (0.025mmol) (4), 0.176g (1mmol) hexamethylene diamine, and 235µL diisopropylcarbodiimide (DIC) (1.5mmol) (Avocado) were stirred under N_2 in 35mL CH_2Cl_2/ acetonitrile (1:1 v/v) for three days. SMM chains precipitated from solution. Excess DIC, diamine, and unchained Mn12-acetate were removed by decantation. 35mL acetonitrile was added and the solution was decanted again. The sample (7) was dried under vacuum.
3.2 – Results and Discussion

3.2.1 – Peptide/nanoparticle assemblies

Figure 3.5a shows the CD spectra of unlabelled peptide (1) (black squares), cobalt-peptide (2) (red circles), and SMM-peptide (3) (green triangles); figure 3.5b shows the literature reported spectra of poly-L-lysine as a reference for the trends in helix-coil transitions. The typical “camel’s hump” between 215-230nm that is characteristic of α-helices are present in all three. The spectrum of (1) appears to be that of a combination of a random-coil and α-helix, probably because of the short length of the peptide and low concentration. The spectrum of (3) has significant contributions from the Mn12-tBu/NO₃, especially in the carboxylate region around 280nm.

![Figure 3.5](image.png)

**Figure 3.5** a) Circular dichroism spectra of peptide (1) (black square), cobalt-peptide (2) (red circle), and SMM-peptide (3) (green triangle). b) Spectra of poly-L-lysine, taken from [Campbell84].

While single-molecule magnets because of their size are the ideal superparamagnets for inducing conformational changes in protein secondary structures, the presence of the magnetic field changes the CD spectra of a single-molecule magnet, indicating a coupling between the induced magnetic dipole (spin) and the absorption of circularly polarized light. Future work should focus greatly on this coupling. However, at this time, not enough data on the field-induced changes in the CD spectra of SMMs is available to
be able to report the effect of the magnetic field on the peptide conformation, since the contributions of the SMM to the spectrum of (3) in the presence of a magnetic field cannot be subtracted out with the available data.

3.2.2 – Field induced changes in secondary protein structures

Figure 3.6a shows the effect of the magnetic field on the spectrum of (2) with (black) and without (red) the magnetic field. The magnetic field appears to induce random coil formation in (2) (black), as evident by the greater (linear) slope in the curve at lower wavelengths, between 210-220nm, at which the contributions of the random-coil to the spectra should dominate. Given the space constraints inside the spectrometer, only a very small permanent magnet was placed under the cuvette, so it is likely that only a small fraction of the nanoparticles were magnetized. The conformational changes should be far more detectable with a different experimental setup in which the entire cuvette is surrounded by magnets. These efforts are currently underway, as well as larger scale preparations of (2) for less noisy spectra, extensive temperature scans to evaluate the effect of cobalt nanoparticles on the stability of the α-helix, and the preparation of assemblies with different peptides that may form more stable α-helices.

Figure 3.6 Induced α-helix to random-coil transition due to induced dipole interactions between cobalt nanoparticles. a) Circular dichroism spectra with (red) and without (black) a small magnetic field. b) Spectra of poly-L-lysine. Image from [Campbell84].
3.2.3 – Synthesis of molecular magnets and magnetic wires (7)

While site-specific ligand abstraction in SMMs has been shown in the literature [Christou01], reactions attacking those chemically labile sites have yet to be shown. Figure 3.7 shows the FTIR spectra of Mn12-acetate (4) and the molecular magnetic wires (7). It is believed that the SMM chains (7) were formed via deprotonated amide bonds at the chemically accessible carboxylate sites in which both oxygen and nitrogen are involved in coordinating the ions. It is important to note that these changes in the spectra persist after multiple purification steps, which would indicate that they are not from unreacted hexamethylene diamine.

![Figure 3.7 FTIR spectra of Mn12-acetate (1) and Mn12-acetate molecular wires (8).](image)

The spectrum for (7) shows an increase in intensity in the alkyl regions (2929 and 2857cm\(^{-1}\)) relative to carboxylate peaks (1567 and 1417cm\(^{-1}\)) and a new peak at 1468cm\(^{-1}\) (CH\(_2\) bend) that indicates the presence of the methylene groups from hexamethylene diamine. The significant increase in relative intensity of the peak at 1567cm\(^{-1}\) versus 1417cm\(^{-1}\) indicates the presence of N-H groups (N-H bends). Changes in the carboxylate region of (7) (broadening, increase in the 1567cm\(^{-1}\) peak intensity, and the new peak at 1468cm\(^{-1}\)) may also be attributable to the formation of deprotonated amides that replace four (25%) of the carboxylate ligands. The most telling evidence for the formation of deprotonated amides is the broadening in the \(\nu < 1336\)cm\(^{-1}\) region and
the hump at 1280cm$^{-1}$ in (7), which are unique to metal-chelated nitrogen atoms and are not present in amines [Nakamoto68].

Figure 3.8 a) Proposed route towards molecular wires composed of a string of individual single-molecule magnets with aligned crystalline axes. b) TEM micrograph of molecular wires (7). Scale bar = 10nm.

Figure 3.8 shows a TEM micrograph of (7) which appears crystalline, whereas micrographs of (4) only showed amorphous material. Because the bridging between SMMs occurs along the Jahn-Teller axis, the bridging diamide chemically orders the sample, making it effectively “crystalline.” The diameter of the wire is 0.9nm, which
agrees with the calculated MnO core size from XRD data of (4) [Lis80]. The ordering of wires into arrays is likely due to strong Van der Wals attraction between the wires as the solvent evaporates. EDAX data shows that the material is MnO and not crystalline impurity. The sample did not vaporize under the TEM, ruling out the possibility that the material was unpurified salt from the synthesis ([KMnO₄] and [Mn(CH₃O₂)•4H₂O]).

The TEM micrograph of (7) also shows that the wires do not branch to form Y junctions, indicating that each SMM is bound to only one SMM on each side, even though there are two possible bridging sites at each end of one SMM. This can be explained either by steric hindrance, or by the fact that after one bond is formed between two SMMs, it is far more likely that a second bond will form between the two (by searching the conformational space) before another SMM links to the growing chain (i.e. the time scale of conformational search is shorter than the time scale of diffusion). The straightness of the wires would indicate that each SMM is bridged by two amides to each adjacent SMM, thereby limiting the conformational freedom.

Figure 3.9 M-H curves of Mn12-acetate molecular wires (7) showing room temperature hysteresis.

Figure 3.9 shows a room temperature M-H curve of the wires with no field-alignment. A small amount of hysteresis ($H_c = 72$Oe) can be observed in figure 3.9, despite the fact that (4) is superparamagnetic. The hysteresis loop is narrower than the loop observed in field-aligned (4) ($H_c = 202$Oe) (figure 3.2) because the SMMs in the chain are separated
by hexamethylene (C6) groups that decrease the (distance) dependent spin-interactions between the SMMs in the chain. M-H curves SMMs reacted with ethylene diamine (C2) and 1,12-aminododecane (C12) show no hysteresis. SMMs reacted with ethylene diamine are likely not bridged because the linker molecule is too short, and a wire cannot form. On the other hand, the C12 group is perhaps too long for significant spin interaction to occur between SMMs at room temperature.

The data above indicates the synthesis of a magnetic nanowire that is exactly one molecule wide and shows room temperature hysteresis. In the context of controlling biological molecules, the ability to bridge superparamagnets may allow a protein conformation to be altered, and permanently “turned off” by covalent locking. Observable hysteresis measurements would perhaps allow protein conformational changes to be monitored in vivo using a SQUID magnetometer.
Chapter 4: Wavelength Selective Photochemistry

4.0 – Introduction to wavelength selective photochemistry

This chapter proposes the design of a molecular assembly line based on wavelength selective photochemistry, in which the shuttle is covalently handed-off between monomer units much like ketides in a polyketide synthase. The basis of the system is the ability to use wavelength selective Norrish Type II photocleavage reactions of carbonyls to control the equilibrium of a Michael reaction (section 4.1). Experimental data on the ability to control this equilibrium will be shown, and potential backbones are proposed.

4.0.1 – Light-driven and programmable chemistries

Photochemical reactions are of great importance in organic, bioorganic, and polymer chemistries. Typically, UV irradiation is used as a means for polymerization initiation, intramolecular rearrangements, or photocleavage of protecting groups or traceless linkers [Bochet02] [Kopecky92]. Largely driven by lab-on-chip gene arrays and solid-phase oligonucleotide synthesis, great efforts have been directed towards designing wavelength selective photocleavable chemistries for orthogonal protecting groups (figure 4.1) [Bochet02]. Although it is normally not a concern in protecting group chemistry, photocleavage reactions typically do not yield products that can be reacted to give back the initial molecule. This greatly limits the types of photochemical reactions that can be employed to build a molecular assembly line, which requires reversibility.

Figure 4.1 The role of photocleavable protecting groups in solid-phase DNA synthesis. The blue rectangles represent 5’ photocleavable protecting groups necessary for making gene arrays on a chip by photolithography. Image from http://www.affymetrix.com.
However, such systems do exist, and have been demonstrated in the literature. In an extremely clever route towards reversible DNA ligation, Saito et al. ligated two DNA strands with a cyclobutane group formed by a light-driven 2+2 reaction with 5-vinyl-deoxyuridine (molecule 4 in figure 4.2a) at 366nm. Irradiating the DNA with 300nm light breaks the cyclobutane group and reverses the ligation step [Saito00]. Also, a class of charge-transfer complexes, called TCNQs (figure 4.2b), contains a large number of photochromic zwitterions with highly resolvable absorptions bands [Petty95]. These compounds could be used to build an assembly line based on electrostatic interactions between TCNQs and a charged shuttle.

![Figure 4.2 a] Reversible DNA ligation by light-driven “2+2” addition / photocleavage on a DNA backbone. Image from [Saito00]. b) Structure of a TCNQ photochromic zwitterion.

In truth, there are many routes towards building light-driven molecular machines and electronics (such as cis-trans isomerizations in azo-compounds and Förster resonance energy transfer, or FRET), but ultimately, the above-mentioned chemistries were not explored because of concerns over wavelength selectivity, or the fact that the shuttle/monomer interactions would not be covalent, since our goal here is to construct a purely synthetic analog of polyketide synthase that is capable of covalent hand-off.

### 4.0.2 – Molecular assembly line based on photochemistry

The reaction of choice for this embodiment of the molecular assembly line is the Norrish Type II photocleavage reaction of carbonyl compounds [Norrish38] [Norrish52]; the
Norrish Type II reaction results in an alkene and a terminal ketone or amide. Depending on their constituents, the photocleavage products may also be potential reactants for a base-catalyzed Michael reaction (figure 4.3a) [March92]. Therefore, the relative equilibrium of a Michael reaction can be controlled by UV light. Furthermore, the wavelengths at which the molecules (1-3) cleave are determined by and can be tuned by the R group. An alkyl ketone will cleave at $\lambda = 280\text{nm}$, whereas an aryl ketone and alkyl amide cleave at $\lambda = 310\text{nm}$ and $\lambda = 210\text{nm}$, respectively. It is conceivable that a molecular assembly line could be constructed using this type of competitive Norrish/Michael reaction, as shown in figure 4.3b, where the Michael acceptor (4) is the shuttle and the Michael donor is the antenna. The experimental work in this chapter includes the photocleavage of an alkyl ketone and amide.

\begin{align*}
\text{a)} \\
\begin{tikzpicture}[baseline=(current  bounding  box.center)]
\node[align=center] (a) at (0,0) {\text{hv}}; \\
\node[align=center] (b) at (0,-0.5) {\text{Base}}; \\
\node[align=center] (c) at (4,0) {\text{CN}}; \\
\node[align=center] (d) at (4,-0.5) {\text{O}}; \\
\node[align=center] (e) at (1,0) {\text{CN}}; \\
\node[align=center] (f) at (1,-0.5) {\text{O}}; \\
\node[align=center] (g) at (2,0) {\text{R}}; \\
\node[align=center] (h) at (2,-0.5) {\text{CN}}; \\
\node[align=center] (i) at (3,0) {\text{CN}}; \\
\node[align=center] (j) at (3,-0.5) {\text{O}}; \\
\node[align=center] (k) at (1.5,0) {\text{280nm}}; \\
\node[align=center] (l) at (2.5,0) {\text{310nm}}; \\
\node[align=center] (m) at (3.5,0) {\text{210nm}}; \\
\end{tikzpicture}

\text{where,} \\
(1) \ R = \text{alkyl} \ (280\text{nm}) \\
(2) \ R = \text{aromatic} \ (310\text{nm}) \\
(3) \ R = \text{NR}_2 \ (210\text{nm})
\end{align*}

\begin{align*}
\text{b)} \\
\begin{tikzpicture}[baseline=(current  bounding  box.center)]
\node[align=center] (a) at (0,0) {\text{NC}}; \\
\node[align=center] (b) at (0,-0.5) {\text{O}}; \\
\node[align=center] (c) at (2,0) {\text{CN}}; \\
\node[align=center] (d) at (2,-0.5) {\text{O}}; \\
\node[align=center] (e) at (4,0) {\text{NC}}; \\
\node[align=center] (f) at (4,-0.5) {\text{O}}; \\
\node[align=center] (g) at (1,0) {\text{CN}}; \\
\node[align=center] (h) at (1,-0.5) {\text{O}}; \\
\node[align=center] (i) at (3,0) {\text{CN}}; \\
\node[align=center] (j) at (3,-0.5) {\text{O}}; \\
\node[align=center] (k) at (1.5,0) {\text{280nm}}; \\
\node[align=center] (l) at (2.5,0) {\text{310nm}}; \\
\node[align=center] (m) at (3.5,0) {\text{210nm}}; \\
\end{tikzpicture}

\begin{tikzpicture}[baseline=(current  bounding  box.center)]
\node[align=center] (a) at (0,0) {\text{NC}}; \\
\node[align=center] (b) at (0,-0.5) {\text{O}}; \\
\node[align=center] (c) at (2,0) {\text{CN}}; \\
\node[align=center] (d) at (2,-0.5) {\text{O}}; \\
\node[align=center] (e) at (4,0) {\text{NC}}; \\
\node[align=center] (f) at (4,-0.5) {\text{O}}; \\
\node[align=center] (g) at (1,0) {\text{CN}}; \\
\node[align=center] (h) at (1,-0.5) {\text{O}}; \\
\node[align=center] (i) at (3,0) {\text{CN}}; \\
\node[align=center] (j) at (3,-0.5) {\text{O}}; \\
\node[align=center] (k) at (1.5,0) {\text{280nm}}; \\
\node[align=center] (l) at (2.5,0) {\text{310nm}}; \\
\node[align=center] (m) at (3.5,0) {\text{210nm}}; \\
\end{tikzpicture}

\begin{tikzpicture}[baseline=(current  bounding  box.center)]
\node[align=center] (a) at (0,0) {\text{NC}}; \\
\node[align=center] (b) at (0,-0.5) {\text{O}}; \\
\node[align=center] (c) at (2,0) {\text{CN}}; \\
\node[align=center] (d) at (2,-0.5) {\text{O}}; \\
\node[align=center] (e) at (4,0) {\text{NC}}; \\
\node[align=center] (f) at (4,-0.5) {\text{O}}; \\
\node[align=center] (g) at (1,0) {\text{CN}}; \\
\node[align=center] (h) at (1,-0.5) {\text{O}}; \\
\node[align=center] (i) at (3,0) {\text{CN}}; \\
\node[align=center] (j) at (3,-0.5) {\text{O}}; \\
\node[align=center] (k) at (1.5,0) {\text{280nm}}; \\
\node[align=center] (l) at (2.5,0) {\text{310nm}}; \\
\node[align=center] (m) at (3.5,0) {\text{210nm}}; \\
\end{tikzpicture}

\text{Figure 4.3 a) General reaction scheme for a Michael reaction in which the reaction equilibrium is driven by a competitive Norrish Type II photocleavage reaction. b) The repeating (1-2-3) unit of a molecular assembly line based on wavelength selective photocleavage of carbonyl compounds.}
The general molecule was designed as it was for several reasons. First, the methyl groups at the \(\gamma\)-carbon sterically promote the Norrish Type II pathway for the photocleavage reaction, which could otherwise lead to the formation of an undesirable cyclobutane compound [Wagner83] by a Norrish Type I pathway. The p-tolyl group also favors the reaction taking a Norrish Type II pathway. Although it lowers the probability that the photocleavage reaction follows a Norrish Type II pathway versus a Norrish Type I pathway, the nitrile group makes (4) a far better electrophile and Michael acceptor.

It should be noted that the photocleavage of the o-tolyl version of (1) has been shown to be quantitative and yielded only the Norrish Type II products [Kim97]. The p-tolyl version is not as efficient because of the larger distance between the 1 and 4 carbons in the bi-radical intermediate, but as will be discussed in section 4.2, the reaction still only gave Norrish Type II products. The p-tolyl group was chosen over the o-tolyl group because of the availability of the reagent. A p-tolyl group was used instead of a benzene group to conserve the directing effects of having a substitution that would be required for linking the molecule to a solid-support.

One drawback of building a supramolecular molecular assembly line is the difficulty in synthesizing a linear backbone with reactive functionalities at exact, controlled locations... yet another reason to marvel at linear biological systems like poly-peptides and oligonucleotides. The most promising route is likely a ring-opening metathesis polymerization (ROMP), which is the route used in synthesizing the “endless” cyclic polymers reported by Grubbs et al. [Grubbs02] and the “universal polymer backbone” by Weck et al. [Weck03]. Many researchers have used DNA as a backbone for pre-programmed chemical reactions [Liu01] [Sheppard01], but the destructive nature of UV irradiation at wavelengths required for carbonyl photocleavage reactions makes DNA incompatible with the proposed chemistry.

Despite the difficulties in creating a backbone and limited applications in assembly given the destructive nature of UV irradiation, pursuing this avenue is extremely interesting for various reasons. A system with the ability to covalently hand-off a molecule would be a completely synthetic analog of the polyketide synthase. Wavelength selective photocleavable groups are extremely powerful as protecting groups. Furthermore, the
Michael reaction is a very general and useful reaction in organic chemistry, and the ability to drive the equilibrium of certain Michael reactions may prove useful, especially as the notions of *programmable chemistry* and *dynamic covalent chemistry* [Lehn99] [Stoddart02] become more and more prevalent.
4.1 – Experimental

All reagents were of the highest purity available and were used as received from Sigma-Aldrich unless otherwise indicated. Reactions under nitrogen were performed using standard techniques. $^1$H-NMR spectra were taken on a Varian Unity 300MHz NMR spectrometer in CDCl$_3$. FTIR was taken using a Digi-Lab Excalibur FTS 3000 (neat, 64 scans, 4cm$^{-1}$ resolution). HPLC chromatographs were obtained on a Varian Pro Star HPLC (monitored by UV absorption at 254nm).

Photocleavage reactions were performed under nitrogen in a photochemical micro-reaction assembly (Ace Glass model 7880) in cyclohexane (1mM) using a Pen-Rey 5.5W low-pressure mercury UV-lamp. (1) was irradiated with $\lambda > 290$nm by filtering lower wavelengths with borosilicate glass. (3) was irradiated with $\lambda \sim 215$nm by filtering with a 3 : 1 : 0.5 (v:v) mixture of ethanol : water : acetone with constant bubbling of nitrogen through the solution [Muel69]. HPLC of (1) and (3) were performed on a normal-phase silica column with 70:30 and 50:50 (v:v) heptane/THF as eluent, respectively.

The synthetic schemes for (1) and (3) are provided in figure 4.4.

![Figure 4.4](image-url)

**Figure 4.4** a) Synthetic scheme for 5-(p-tolyl)-5-cyano-4,4-dimethyl-2-pentanone (1). b) Synthetic scheme for 4-(p-tolyl)-4-cyano-3,3-dimethyl-N,N-dimethylbutaneamide (3).
**Synthesis of 5-(p-tolyl)-5-cyano-4,4-dimethyl-2-pentanone (1)**

660µL p-tolyl-acetonitrile (5mmol) and 572µL mesityl oxide were stirred in 50mL THF/HMPA (v:v 80:20) for 30 minutes at -78°C under nitrogen. 5mL of 1M LiHMDS in THF (5mmol) was added dropwise. The reaction was brought slowly to room temperature over 90 minutes, and stirred for two days at room temperature. The solution was extracted using ether / 1M HCl, and purified by silica gel flash chromatography (20:80 ethyl acetate/hexanes, v:v). After solvent removal by rotary evaporation, the sample was dried under high vacuum overnight. \(^1\)H-NMR \(\delta\) (ppm): 7.45-7.25 (4H, m), 4.82 (1H, s), 2.98 (3H, s), 2.36 (3H, s), 2.44-2.11 (2H, AB quartet), 1.25 (3H, s), 1.08 (3H, s). FTIR 2236 (m, s, CN), 1712.1 (s, br, CO).

**Synthesis of 4-(p-tolyl)-4-cyano-3,3-dimethyl-N,N-dimethylbutaneamide (3)**

557µL 3,3-dimethylacryloyl chloride (5mmol) and 6.25mL 2M dimethylamine in THF (2.5x, 12.5mmol) were stirred under nitrogen at 0°C for 30 minutes in 43.25mL CHCl\(_2\). The compound was purified by extraction with ether / 1M HCl. Ether was removed by rotary evaporation to yield 3-methyl-N,N-dimethyl-2-buteneamide.

5mL of 1M LiHMDS in THF (5mmol) was added dropwise to a solution of 660µL p-tolyl-acetonitrile (5mmol) in 25mL THF/HMPA (80:20, v:v), and stirred for 30 minutes at -78°C under nitrogen. 25mL of 0.2M 3-methyl-N,N-dimethyl-2-buteneamide in THF/HMPA was added dropwise. The reaction was brought slowly to room temperature over 90 minutes, and stirred for two days at room temperature. The solution was extracted using ether / 1M HCl, and purified by silica gel flash chromatography (70:30 ethyl acetate/hexanes, v:v). After solvent removal by rotary evaporation, the sample was dried under high vacuum overnight. \(^1\)H-NMR: \(\delta\) 7.26-7.12 (4H, m), 4.82 (1H, s), 2.97 (6H, s), 2.34 (3H, s), 2.43-2.11 (2H, AB quartet), 1.23 (3H, s), 1.05 (3H, s). FTIR (cm\(^{-1}\)) 2235 (m, s, CN), 1643 (s, br, CONR\(_2\)).
4.2 – Results and Discussion

Figure 4.5a and 4.5b show the respective HPLC chromatographs of (1) and (3) after 48 hours UV irradiation at $\lambda > 290$nm and $\lambda = 215$nm, respectively. In each chromatograph, the left peak represents the cleavage product / Michael acceptor, 3-(p-tolyl)-3-cyano-2-methyl-2-butene (4), and the right peak represents the initial molecules (1) and (3) / Michael products. The Michael donors, acetone and N,N-dimethylacetamide, are not present in the spectra, as they are likely removed by water / heptane extraction during HPLC sample preparation. As expected, the cleavage reactions are wavelength selective as determined by thin-layer chromatography (TLC) analysis. Also as expected, the cleavage reaction followed a Norrish Type II pathway, and not a Type I cyclization, as determined by NMR analysis of products purified by silica-gel flash chromatography - (4) $^1$H-NMR $\delta$ (ppm): 7.33-7.17 (4H, m), 2.38 (3H, s), 2.25 (3H, s), 1.92 (3H, s).

Figure 4.5 a) and b) HPLC chromatographs of (1) and (3) after 48 hours UV irradiation, respectively.

After 48 hours, the sample was aliquoted into two samples by removing half the sample volume from each reaction mixture, and an equimolar amount of LiHMDS was added to each sample. One aliquot was allowed to react in the presence of UV irradiation, and another was stirred under nitrogen in the dark at room temperature. After 48 hours, the relative equilibrium of the Michael reaction between (4) and the acetone enolate or the N,N-dimethylacetamide amidine was controlled by the UV irradiation, as shown in figure 4.6.
The basic principle of influencing the Michael reaction equilibrium has been proven, but the current experimental methods are limited in their ability to achieve quantitative yields of any product in the simultaneous presence of UV and lithium base. It is also impossible to make quantitative kinetic and thermodynamic comparisons between the compounds in the simultaneous presence of UV and lithium base. First, since the emission of the lamp is centered about 254 nm, the number of photons in the experiment
is extremely low after wavelength filtering. Furthermore, the number of photons per unit time between the photocleavage reactions of (1) and (3) are not equal, making it difficult to compare the two. Second, in the case of compound (1), the lithium base catalyzes the Aldol condensation of acetone, forming mesityl oxide; this prevents the Michael reaction from ever being quantitative. It could only be driven to quantitative yields in the presence of an excess of LiHMDS and acetone, and in the absence of UV irradiation. In the case of (3), the N,N-dimethylacetamide amidine is a fairly poor nucleophile in comparison to an acetone enolate, changing the reaction kinetics between the two systems.

Figure 4.7 a) An intramolecular toggle switch driven by UV light. b) Potential synthetic scheme towards (5).
Therefore, future experiments must first employ single-wavelength UV lasers with known powers or photon count per unit time. Second, the reactive functionalities – Michael acceptor, enolate, and amidine – must all be included in the same molecule so that the experimental time-scales are limited by the time-scale of bond-rotation, and not diffusion. This would prevent the formation of side-products like mesityl oxide and allow kinetic constants to be more easily determined. Figure 4.7 proposes a potential molecule to be synthesized in the future, an intramolecular toggle-switch (5); a potential synthetic scheme towards (5) [Mosley, UPR] is also given. The methyl groups that provide steric hindrance and the aromatic group that promotes hydrogen abstraction adjacent to the nitrile are not required. 3D-modeling (ChemDraw 3D Ultra, Cache) showed that products of the photoreaction following a Norrish Type I path, or a path via 1-5, 1-6, or 1-7 bi-radical intermediates, are unfavorable because of great ring strain.
Chapter 5: Summary and further work

While the “molecular assembly line” is, at this point, only a generalized scheme, the experimental results in this thesis have shown that its construction is within reason, as the basic requirement of the system as described in Chapter 1 - wavelength (or frequency) selective control over the interactions between monomeric antennae and a shuttle – has been met.

In the first approach, “RF Biology” (Chapter 2), an oligonucleotide was selectively dehybridized by coupling energy from 300MHz RFMF into covalently-attached 2.8nm cobalt nanoparticle antennae. In the future, the nanoparticle size distribution and the coupling chemistry to the oligonucleotides must be improved, and fluorescence measurements must be made with an internal thermometry standard in solution. Furthermore, a localized heating model has been proposed for the selective dehybridization of the oligonucleotide (as opposed to some mechanical resonance effect), but this has yet to be experimentally verified.

In the second approach, “Wavelength-selective photocleavage of carbonyl compounds” (Chapter 4), the equilibrium of a Michael reaction between an alkene acceptor and enolate or amidine donor was driven by UV irradiation at the absorption wavelength of the carbonyl bond in the resultant Michael product. This approach is particularly interesting because it holds the possibility of constructing a purely synthetic analog to the polyketide synthase in its ability to covalently “hand-off” small molecules. In the future, experiments must employ single-wavelength lasers, and a molecule must be synthesized that contains an alkene acceptor and both an enolate and amidine donor.

Upon the synthesis of a polymerized molecular assembly line, the issue of polymer chain coiling will have to be resolved. For example, the persistence length of double-stranded DNA is approximately 50nm [Bustamente00], and therefore, the length of a molecular assembly line built out of RF Biology parts will be limited to 50nm without employing external forces, like electric fields to align the phosphate backbone.

The experimental work in this thesis represent the most preliminary experimental steps towards constructing a molecular assembly line. In truth, the results are simple “proofs-
Summary and future work

of-concept” that interactions between molecular components can be driven by external inputs, like RFMF and UV light, with wavelength/frequency selectivity. Until the synthesis of a fully polymerized molecular assembly line, the question will still remain whether the molecular assembly line is a useful construction, or whether its utility is merely as a “proof of concept.”

Perhaps the most interesting outcomes will be the applications of the chemistries developed during the process but are outside the realm of building a molecular assembly line. For instance, the goal of controlling biological molecules so that they could be used as components in a molecular assembly line spawned the synthesis of molecular wires in Chapter 3 of this thesis. The ability to selectively break hydrogen bonds has great potential applications in biophysics and bioengineering, and the wavelength selective photocleavage reactions might have great potential applications in protecting group chemistry.

Only further experimental work can answer these questions, but given the initial set of results here, the outlook is promising.
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***

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***

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From left to right: (bottom) Peter, Joe, and Kie-Moon (middle) Vikas, Eric, Will, and Brian (top) Dave M., Dave K., and Saul (not shown): Kim, Vikrant, Sawyer, Brent, Mike, and Murray.
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