Linking Wheels for use in Quantum Information Processing

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Abstract

Heterometallic Cr\textsubscript{7}Ni-containing wheels have been identified as potential qubits for use in quantum information processing. The work described in this thesis details attempts to form a variety of multi-qubit systems for the purposes of better understanding the interactions occurring between the wheels and also forming potential 2-qubit quantum gates using redox/photo-active links.

Mono-substituted wheels of type [Pr\textsubscript{2}NH\textsubscript{2}][Cr\textsubscript{7}NiF\textsubscript{8}(O\textsubscript{2}CCMe\textsubscript{3})\textsubscript{15}(L)], where L is a carboxylate with a pendant coordinating group, have been synthesised for use as bulky ligands in coordination chemistry with metal complexes. Various carboxylates have been substituted into the wheels and the products reacted with first row transition metal complexes in order to extend the series of linked-wheel systems. Many of these novel complexes have been characterised by X-ray crystallography, and in certain cases EPR studies have been undertaken to probe the strength of interactions occurring via different bridging units.

The first well-established substituted wheel, [Pr\textsubscript{2}NH\textsubscript{2}][Cr\textsubscript{7}NiF\textsubscript{8}(O\textsubscript{2}CCMe\textsubscript{3})\textsubscript{15}(O\textsubscript{2}CPy)] (Py = pyridine), has also been used in reactions with second and third row transition metal centres to show its ability to act as a ligand under more harsh conditions. In addition, the disubstituted product [Pr\textsubscript{2}NH\textsubscript{2}][Cr\textsubscript{7}NiF\textsubscript{8}(O\textsubscript{2}CCMe\textsubscript{3})\textsubscript{14}(O\textsubscript{2}CPy)\textsubscript{2}] has been reacted with a copper complex in order to form a polymeric structure.

Purple wheels of type [Cr\textsubscript{7}NiF\textsubscript{3}(O\textsubscript{2}CCMe\textsubscript{3})\textsubscript{15}(EtGu)(H\textsubscript{2}O)] have been linked through a variety of extended organic molecules containing a minimum of two pendant pyridyl groups. Use of a ligand containing four pyridyl groups, 2,2′:4,4′:4″,4″″-quaterpyridyl (qpy) produced a three-wheel-containing system, creating an interesting modification of the wheel backbone not seen before.

A series of transition metal complexes, containing qpy and its extended derivative bbpe, has been synthesised in order to form linked-wheel systems. While none of these systems has to date yielded a crystal structure, a significant amount of evidence has been collected to confirm successful formation of the desired products. UV-vis spectroscopic and electrochemical measurements show that these compounds are photo and/or redox-active, and preliminary luminescence studies indicate that the presence of the wheels quenches the emission from metal-to-ligand charge-transfer excited states.

A series of mixed-wheel systems has been synthesised by reacting four different mono-substituted green wheels with four purple wheels (Cr\textsubscript{7}M, where M = Ni, Co, Mn or Zn). A total of fourteen novel systems has been formed, with two of the attempted reactions failing to occur. EPR studies on Cr\textsubscript{7}Ni-Cr\textsubscript{7}Ni linked systems show that the strongest interaction occurs when using isonicotinate as a link, with 4-pyridazincarboxylate giving the weakest coupling. Studies have also been undertaken on Cr\textsubscript{7}Ni-Cr\textsubscript{7}Mn and Cr\textsubscript{7}Ni-Cr\textsubscript{7}Zn systems, with the former showing interactions and the latter showing none.

Initial work to link wheels together indirectly via their templates was ultimately unsuccessful, but a variety of potential linker molecules were synthesised and are described.
Declaration

I hereby declare that no portion of the work referred to in this thesis has been submitted in support of an application for another degree or qualification of this or any other university or other institute of learning.

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Abbreviations

2,2´-bpy 2,2´-bipyridyl
4,4´-bpy 4,4´-bipyridyl
4-pic 4-picoline
4-pyac 4-pyridylacrylate
4-pyd 4-pyridazine carboxylate
BOC tert-butoxycarbonyl
bbpe 4,4´-bis-[(E)-2-(4-pyridyl)vinyl]-2,2´-bipyridyl
bpe (E,1,2-bis(4-pyridyl)ethylene
bph (E,E,E)-1,6-bis(4-pyridyl)-1,3,5-hexatriene
bpvb 1,4-bis[E-2-(4-pyridyl-ethenyl]benzene
bpym 2,2´-bipyrimidine
CV cyclic voltammetry
cyclen 1,4,7,10-tetraazacyclododecane
DCM dichloromethane
DIC diisopropylcarbodiimide
DMF N,N-dimethylformamide
DMSO dimethylsulfoxide
EPR electron paramagnetic resonance
ESI-TOF electrospray ionisation time-of-flight
ES-MS electrospray mass spectrometry
Et ethyl
EtGlu N-ethyl-D-glutamine
Et2O diethyl ether
EtOH ethanol
fac facial
H-bond hydrogen bond
hfac hexafluorooxycetacetate
HOBT 1-hydroxybenzotriazole
HOPG highly oriented pyrolytic graphite
IR infrared
ISNA isonicotinate
\( \lambda_{\text{max}} \) wavelength of maximal absorption
MALDI-MS  matrix assisted laser desorption ionisation mass spectrometry
Me  methyl
MeCN  acetonitrile
MeOH  methanol
mer  meridional
MLCT  metal to ligand charge transfer
MS  mass spectrometry
NMR  nuclear magnetic resonance
Pr  propyl
py  pyridine
pytpy  4´-(4´´-pyridyl)-2,2´:6´:2´´-terpyridine
QIP  quantum information processing
qpy  2,2´:4,4´´:4´,4´´´-quaterpyridyl
qubit  quantum bit
SAM  self assembled monolayer
SMM  single molecule magnet
sol  solvent
SQUID  superconducting quantum interference device
STM  scanning tunnel microscopy
tacen  1,4,7-triazacyclononane
'tBu  tert-butyl
TFA  trifluoroacetic acid
THF  tetrahydrofuran
TLC  thin layer chromatography
UV  ultraviolet
VdW  Van der Waal’s forces
vis  visible
VT  variable temperature
XPS  x-ray photoemission spectroscopy
Dedicated to my granddad, William Chatfield, my first great educator.
Chapter 1 - Introduction
1.1 – OVERVIEW

Since the design of “Baby”, the first computer as we know it, in Manchester in 1948\(^1\) the field of computer technology has been continuously advancing with the speed and memory capacity of new computers improving greatly year by year. The rate of advancement is slowing however, primarily due to a limit on the miniaturisation scale of silicon chips.\(^2\) Consequently demand has increased for the development of a new genre of computing, and since the 1970s the concept of quantum computing has been increasingly studied. Theoreticians have long been designing concepts by which quantum computing could become a reality, in a top-down approach, and more recently chemists have become involved trying to solve the problem from a bottom-up approach.

Recent theoretical studies by Loss and Meier have suggested that multi-spin clusters may be suitable for use as qubits, data storage units in quantum information processing, as long as their overall molecular spin is a stable two state system.\(^3,4\) Consequently, heterometallic Cr\(_7\)Ni-containing wheels synthesised by the Winpenny group, which have a ground state spin of \(\frac{1}{2}\), show promise for this application.\(^5,6\) As in regular computers, quantum computers require logic gates containing multiple qubits. These must be linked together through a molecule that allows a selective inter-cluster exchange enabling entanglement and thus communication between the qubits.

A range of synthetic approaches to link magnetic molecular wheels are considered here, both using the templates of these wheels and also directly through their backbones. Numerous compounds are considered as linkers, some for the purpose of observing permanent effects of communication on the wheels, and others with the aim of creating a molecular switch. Focus on switchable linkers is primarily on the use of photo- and/or redox-active molecules, such as viologens or metal-to-ligand charge-transfer (MLCT)-active transition metal complexes. These have been chosen as it is expected that they will have sufficiently long excited-state lifetimes to affect communication between the wheels.
1.2 PARAMAGNETISM

1.2.1 Electron Spin and Magnetic Fields

Species that contain unpaired electrons demonstrate a phenomenon known as paramagnetism. This results in an attraction of the unpaired electrons to a magnetic field, dependent upon the field strength and temperature. Electrons have a property known as spin, with a magnetic quantum number of $M_S = \pm 1/2$. The unpaired electrons can align themselves with or against the magnetic field, with it being much more favourable to align with the field, creating an energy gap between the two spin states. The energy gap grows proportionally to the magnitude of the magnetic field. The effect of the magnetic field on the energy gap, known as the Zeeman splitting, is depicted in Figure 1 and quantified by Equation 1.

**Equation 1:** \[
\Delta E = g\mu_B H
\]

$\Delta E$ = energy gap, $g$ = g-factor, $\mu_B$ = Bohr magneton, $H$ = magnetic field, $S$ = spin state.

![Figure 1. Influence of magnetic field on energy gap between spin states.](image)
The state that aligns with the field is known as the ground state. If the thermal energy is less than the energy gap between the ground state and other excited states \((kT << \mu_B H)\) then the spins will populate the ground state exclusively. If, however, the thermal energy is greater than the energy gap \((kT >> \mu_B H)\) then the spins align themselves randomly.

**Equation 2:**
\[
\chi = \frac{\delta M}{\delta H}
\]

The magnetic susceptibility, \(\chi\), of a substance (Equation 2) is defined as the effect that a magnetic field, \(H\), has on the magnetisation, \(M\). If a substance has a high magnetic polarisation then it can obtain a magnetic field itself thus becoming a magnet. It has been found in certain paramagnetic substances that the magnetic susceptibility is inversely proportional to the temperature, according to Curie’s Law (Equation 3).

**Equation 3:**
\[
\chi = \frac{C}{T}
\]
Where \(C = \frac{N_A g^2 B^2}{4K}\)

\(C =\) Curie constant, \(N_A =\) Avogadro’s constant \((6.023 \times 10^{23} \text{ mol}^{-1})\), \(g =\) electron g factor, \(B =\) Bohr magneton \((9.27 \times 10^{-24} \text{ J.T}^{-1})\), \(K =\) Boltzmann constant \((1.38 \times 10^{-23} \text{ m}^2\text{kg.s}^{-2}\text{k}^{-1})\)

### 1.2.2 Coupling between Magnetic Centres

If there are multiple paramagnetic centres in proximity to one another, then coupling interactions can occur between the unpaired electrons. This can either occur directly, as in bulk metals for example, or through bridging ligands in metal complexes. The unpaired electrons can couple mutually parallel, increasing the size of the overall magnetic dipole in a ferromagnetic interaction. They can also couple antiparallel, cancelling out the dipole in an antiferromagnetic interaction (Figure 2). If the spins couple anti-ferromagnetically but the sizes of the magnetic dipoles differ then a ferrimagnetic interaction is observed, leaving the substance or molecule with an overall magnetic moment.

![Figure 2. Representations of coupling interactions between magnetic centres.](image-url)
If the magnetic coupling is enabled by a bridging ligand then an effect known as superexchange occurs. The magnetic coupling between the spins can be described by the isotropic Heisenberg Hamiltonian, seen in Equation 4, where $J_{i,j}$ is the coupling constant between the spin of $i$, $S_i$, and the spin of $j$, $S_j$. If $J$ is positive, then the coupling between the metals is ferromagnetic, and if it is negative the coupling is anti-ferromagnetic.

$$\hat{H}_{HB} = \sum_{i<j} J_{i,j} S_i \cdot S_j$$

Equation 4:

If the metal centres in a large cluster are coupled ferromagnetically to each other it is possible to obtain clusters with extremely large ground-state spins. The largest overall spin for a single molecule recorded to date being $\frac{83}{2}$ for a Mn$_{19}$ cluster, with 12 Mn$^{III}$ and 7 Mn$^{II}$ ions all being coupled together ferromagnetically.

1.2.3 Single Molecule Magnetism

“Single molecule magnet” (SMM) is a term used to describe a single molecule that can demonstrate magnetic memory behaviour, as opposed to more traditional magnetism that deals with bulk materials. The SMM phenomenon was first demonstrated in 1991 with a Mn$_{12}$-containing molecule, [Mn$_{12}$O$_{12}$(O$_2$CMe)$_{16}$(H$_2$O)$_4$], synthesised by Lis in 1980. This cluster is comprised of eight Mn$^{III}$ ions, each with a spin of 2, and a central cluster of four $S = \frac{3}{2}$ Mn$^{IV}$ ions. The Mn$^{III}$ and Mn$^{IV}$ clusters are anti-ferromagnetically coupled to each other giving the cluster an overall ground-state spin of 10 (Figure 3).

Figure 3. Crystal structure of the first “Mn$_{12}$” cluster and the arrangement of the spins in the ground state cluster.
Having a high spin alone does not ensure SMM behaviour. Another requirement is a large negative magnetic anisotropy (D), which, when coupled with a high overall spin (S), enables a slow relaxation of magnetisation upon application of a magnetic field (Equation 5). Finally, intermolecular interactions must be negligible, so that the magnetic response is of the individual molecule and not of the bulk material.

Equation 5: \[ U = S^2|D| \]

This barrier to relaxation of magnetisation (U) gives rise to a hysteresis loop in a graph of the magnetisation versus the magnetic field (Figure 4), showing that an SMM can retain memory of the external magnetic field. One diagnostic characteristic of the hysteresis loop of an SMM is that the curve is not smooth, but rather descends in "steps" as a result of quantum tunnelling, which occurs when electrons jump to different states, passing barriers they should not have enough energy to overcome.\(^\text{13}\)

![Figure 4. Hysteresis loops of a Mn\(_{12}\) cluster a) varying temperature at a fixed sweep rate and b) varying sweep rate strength at a fixed temperature.\(^\text{13}\)](image)

Studies are ongoing to determine whether modifications can be made to Mn\(_{12}\) SMMs in order to give the maximum possible barrier to magnetisation relaxation.\(^\text{14}\) Most reported SMMs contain Mn, however several others have been reported using other transition metal centres, including Fe(II)\(^\text{15}\), Fe(III)\(^\text{16}\), Co(II)\(^\text{17}\), V(III)\(^\text{18}\) and Mo(III)\(^\text{19}\). Another avenue that is enjoying some success is mixed d-f metal clusters\(^\text{20}\), and recently f-block metal clusters.
have also been shown to be good SMMs, with a Dy(III)$_4$ cluster showing the highest known barrier to relaxation of magnetisation.$^{21}$

1.3 QUANTUM COMPUTING

The concept of quantum computing was suggested as early as 1981 by Richard Feynman, with the idea being that quantum effects would be used as opposed to electronic effects in computing.$^{22}$ However, it was not until Shor$^{23}$ and Grover$^{24}$ published algorithms which showed how the theory could be put into practice, that many different approaches have been considered. Despite these efforts, a system that is functional on a practical scale remains elusive.

1.3.1 Qubits

A quantum bit (qubit) is a unit of quantum information. It is analogous to a bit in a regular data storage system, and thus requires a two-state system (comparable to 1 and 0 of a regular bit). The only difference is that where a regular bit can be $|1\rangle$ or $|0\rangle$, a qubit can be 1, 0 or a superposition of the two. Consequently, any two-state system may be considered as a qubit, and various single-spin candidates have been outlined, such as electron spin and nuclear spin. These are attractive as they are natural two-state systems (spin up and spin down) and because their decoherence times are unusually large for what is expected of their spin degree of freedom.$^{25}$ Another possibility is a multiple state system, in which two states are sufficiently isolable so as to be considered independently of any other excited states.

In 2003, a theoretical publication by Loss $et$ $al.$ outlined the possibility of using multi-nuclear metallic clusters as qubits.$^{3,4}$ The candidates for this were not only 1D, in the forms of odd-numbered spin chains, but also 2D in the shape of bipartite lattices, geometrically frustrated systems such as anti-ferromagnetically coupled triangles and larger molecular clusters such as the heterometallic wheels synthesised by Winpenny $et$ $al.$ Provided that these compounds have a well isolated, single spin ground-state of $S = \frac{1}{2}$, the advantages of using molecules instead of single spins are great as the control over them needs only to be on the molecular scale instead of the atomic scale.
1.3.2 Quantum Gates

In order for qubits to be applied to working computational systems, it is necessary to form a two-qubit quantum gate. This principle was first suggested by Deutsch in 1985\textsuperscript{26}, who compared a quantum logic gate to a conventional logic gate. To realise such a gate, two qubits with $S = \frac{1}{2}$ must be linked together by a molecule that allows a degree of electronic coupling between the qubits, giving an inter-cluster magnetic exchange interaction, $J^*$, that can be modulated by some kind of physical change within the linking unit (Figure 5).

This intramolecular exchange allows entanglement of the qubits, which means that the qubits may no longer be considered as individuals but as one system. By entanglement of an n-qubit system, $2^n$ states are obtainable. QIP algorithms would take advantage of this entanglement to perform certain tasks exponentially quicker than current computer systems.

The linked ring assemblies we aim to synthesise during this project are designed to possess the properties characteristic of such a quantum gate and therefore may prove to be promising prototype systems.

Figure 5. A schematic of a universal set of quantum gates.
1.4 CYCLIC METALLIC CLUSTERS IN THE LITERATURE

Figure 6. A selection of monolayered wheels; Lippard’s “Ferric Wheel”\textsuperscript{27} and Christou’s “Gallic Wheel”.\textsuperscript{28}

The first homometallic wheel published outside of the USSR was Lippard’s “Ferric Wheel” in 1994\textsuperscript{27}, a decametallic cyclic cluster formed by self-assembly from a mixture of $[\text{Fe}_3\text{O(O}_2\text{CCH}_2\text{Cl)}_6(\text{H}_2\text{O})_3]$(\text{NO}_3)\cdot4\text{H}_2\text{O}$, Fe(\text{NO}_3)\cdot9\text{H}_2\text{O}$ and methanol (Figure 6). Each of the ten Fe\textsuperscript{III} ions are bridged by one monochloroacetate and two methoxide ligands, forming an almost planar arrangement of metal centres. Since then, the catalogue of metallic wheels in the literature has been ever expanding, with some of the more impressive cyclic structures including a single-layered Ga\textsubscript{20} “Gallic Wheel”\textsuperscript{28} (Figure 6) and a number of multilayered homo-metallic wheels with 84\textsuperscript{29}, 154\textsuperscript{30} and 176\textsuperscript{31} (the largest to date) metal centres in their backbones (Figure 7).

Wheel-shaped complexes have also been synthesised using Fe(II)\textsuperscript{32}, Cr(III)\textsuperscript{33}, V(III)\textsuperscript{34}, V(IV)\textsuperscript{35}, Hg(II)\textsuperscript{36}, Dy(III)\textsuperscript{37}, Ni(II)\textsuperscript{38}, Cu(II)\textsuperscript{39}, Co(II)\textsuperscript{40} and Ti(IV)\textsuperscript{41} in a variety of even numbered sizes, with 4-20 metal centre containing mono-layered wheels being readily accessible. Very few homometallic odd-numbered metallic wheels have been synthesised, only 5 and 9-membered wheels.\textsuperscript{42,43} In addition to the wheels discussed in Section 1.5, a small number of heterometallic even-numbered examples have been published recently.\textsuperscript{44,45}
The metal centres in these wheels are found in a variety of geometries, and it is possible to impose a great deal of control over wheel size by careful consideration of the reaction conditions, solvent and bridging ligands. Most metal centres in these wheel-shaped structures are coupled together anti-ferromagnetically, but ferromagnetic interactions are observed in a few cases, leading to high ground-state spins and in some cases SMM properties.\textsuperscript{46,47}

1.5 – REVIEW OF THE HETERO METALLIC WHEEL FAMILY

1.5.1 – [Cr\textsubscript{8}F\textsubscript{8}(O\textsubscript{2}CCMe\textsubscript{3})\textsubscript{16}]

The family of heterometallic wheels was derived from the homometallic cluster, [Cr\textsubscript{8}F\textsubscript{8}(O\textsubscript{2}CCMe\textsubscript{3})\textsubscript{16}], which was first synthesised in 1985.\textsuperscript{48} Refluxing CrF\textsubscript{3}•4H\textsubscript{2}O and pivalic acid in DMF for 24 h allowed the self-assembly of this homometallic ring, the structure of which was confirmed by X-ray crystallography and can be seen in Figure 8. Each Cr\textsuperscript{III} ion exhibits a slightly distorted octahedral geometry and is bridged to each adjacent Cr\textsuperscript{III} ion via one fluoride and two pivalate anions. The fluorides point into the centre of the wheel and the pivalate ligands take one of two arrangements; either an
equatorial position in the plane of the ring or an axial position alternating in front and behind the ring.

Figure 8. X-Ray Crystal Structure of [Cr₈F₈(O₂CCMe₃)₁₆].

Situated in the cavity of the wheel in Figure 8 can be seen two acetone molecules, demonstrating the ability of such wheels to act as hosts for small organic molecules. Magnetic studies of this wheel showed that the Cr³⁺ ions are coupled antiferromagnetically to each other, giving an overall ground state spin of 0. The properties of this homometallic wheel have been studied in detail and thus it was chosen as a basis for the formation of heterometallic analogues.

1.5.2 – Heterometallic Wheels

1.5.2.1 – Synthesis

It was found that addition of a divalent metal to the reaction mixture allowed a heterometallic wheel to be formed. By using 2NiCO₃•3Ni(OH)₂•4H₂O it was possible to synthesise an octametallic wheel containing seven Cr³⁺ ions and one Ni²⁺ ion in the backbone. Exchanging a trivalent metal with a divalent metal caused the wheel to become anionic instead of neutral and thus a molecule of protonated dimethylamine, which was formed by acid hydrolysis of the DMF, sits in the cavity of the ring. The homometallic Cr₈ wheel was formed as a by-product that could easily be separated from the anion-cation pair of the heterometallic wheel by column chromatography.
However, the yield of this reaction was only 45% and it was found that a higher yield could be achieved by carrying out the reaction with the carboxylic acid as both solvent and reagent and by use of a primary or secondary amine which, upon protonation, provides a template for the wheel formation. This approach produced a wheel templated about the protonated amine in a relatively high yield of approximately 70% after crystallisation.

1.5.2.2 – Constitution of the Wheels

![Crystal structure of \([\text{Pr}_2\text{NH}_2]\text{[Cr}_7\text{NiF}_8\text{O}_2\text{CMe}_3]\)_{16} and hydrogen bonding in a heterometallic wheel.\(^5\)](image)

In a fashion similar to the \(\text{Cr}_8\) wheel previously described, there are eight metal centres coupled anti-ferromagnetically, seven of which are \(\text{Cr}^{\text{III}}, d^3\) with a spin of \(\frac{3}{2}\) and one of which is \(\text{Ni}^{\text{II}}, d^8\) with a spin of 1. Each metal centre is bridged by two carboxylates and one fluoride, giving a total of sixteen carboxylates and eight fluorides within the backbone. This gives the wheel an overall ground state spin of \(\frac{1}{2}\) and a charge of \(-1\). The wheel is templated about a protonated amine, the amine being held in the cavity of the wheel by hydrogen bonds to three of the fluorides, which point into the centre of the wheel. A molecular structure (Figure 9) helps to visualise this concept.
1.5.2.3 – Physical Properties

This Cr$_7$Ni compound has been well studied in order to verify its potential for use as a qubit in quantum computing. Magnetic studies show that the heterometallic wheel behaves in a similar fashion to the homometallic wheel and the metal centres couple antiferromagnetically to each other, as evidenced by the magnetism, $\chi M T$, of the system decreasing as the temperature is decreased (Figure 10).

![Figure 10. SQUID magnetism studies on a Cr$_7$Ni wheel.](image)

The Q-band EPR spectra of these Cr$_7$Ni wheels confirm that at low temperatures they have a spin of $\frac{1}{2}$, with axial symmetry ($g_{xy} = 1.781$, $g_z = 1.740$) giving rise to a single absorption feature (Figure 11).
1.5.2.4 – Application of the DiVincenzo Criteria to the Wheels

In order for the wheels to be potentially useful in quantum computing, they must follow the four criteria for qubits outlined by DiVincenzo:

i. It must exist in two stable states. For magnetic wheels, this requires that the ground state is a “pure” $S = \frac{1}{2}$ and does not contain a mixture of higher spin states. For Cr$_7$Ni-containing wheels, 99% of the molecules are in a stable $S = \frac{1}{2}$ configuration which means that they can be considered as a two state (binary) system in which $M_S = \pm \frac{1}{2}$ corresponds with the 0 (off) and 1 (on) states.

ii. The energy gap to the first excited state must be large, in order to prevent the population of higher spin states. For Cr$_7$Ni wheels, the energy difference between the $S = \frac{1}{2}$ ground state and the first excited state with $S = \frac{3}{2}$ is 13 K, meaning that at low temperatures $M_S = \pm \frac{1}{2}$ are the only states to be significantly populated.

iii. The $M_S = \pm \frac{1}{2}$ states must separate significantly in energy (due to Zeeman splitting) when an external magnetic field is applied, so that one of them can be selectively populated without significant leakage to higher $S$ states. This creation of a “bit” of information as 0 or 1 is known as initialisation. For Cr$_7$Ni wheels, the Zeeman splitting is 2.4 K (at a field of 2 T) and the energy gap to the lowest level arising from the $S = \frac{3}{2}$
quartet excited state is 9.4 K. Therefore, there should be no leakage and the molecule will still behave as a two state system.

iv. There should be no significant loss of information (decoherence) within the system. For Cr$_7$Ni rings there are two possible sources of decoherence. The first source is a dipolar exchange between molecules, which can be minimised by placing them in a matrix of diamagnetic Cr$_8$ wheels. The second source is hyperfine interactions between unpaired electrons in the metal atoms and nuclear spins in the fluoride ions. These interactions are not strong enough to be resolved in EPR spectroscopy but may still cause decoherence.

1.5.3 – Variations on the Wheels

The majority of this project focuses on the use of Cr$_7$Ni containing wheels, but there is an ever-increasing family of closely related wheels formed by varying one or more features. Changing any or all of the trivalent wheel, the divalent metal, the fluoride, the carboxylate and the ammonium template has led to a family that could number over 1400 rings.

1.5.3.1 – The Trivalent Metal

Chromium is the trivalent metal of choice for this project, but heterometalllic wheels with a M$^{III}$_7Ni backbone have also been synthesised with M = vanadium or iron, as the fluorides of these metals are readily accessible. In air, hydrolysis of iron and oxidation of vanadium can occur, so the synthesis of these wheels must be performed under inert conditions.

More recently, the group 13 metals aluminium, gallium and indium, have also been used in place of chromium. These were used in order to allow study of the paramagnetic M$^{II}$ ion in a wheel arrangement with other diamagnetic ions, and also to test how far this family of rings can be expanded. In all cases, the wheels formed are isostructural with their Cr analogues, but the reaction yields can vary depending on the size of the metal. The best yields of between 70 and 90 % occur with Ga$^{III}$ which is of a similar size to Cr$^{III}$, while the worst (less than 10 %) are for Al$^{III}$, perhaps because its small size is less well-suited for forming an octametallic wheel.
1.5.3.2 The Divalent Metal

Nickel is the divalent metal of choice for this project as when incorporated into a Cr$_7$Ni wheel it gives the most stable molecules and its overall spin of $\frac{1}{2}$ affords the most interesting wheels. However, related wheels have also been formed for Cr$_7$M$^{II}$ where M$^{II} = \text{Co, Fe, Mn, Cd, Zn or Mg}$. All of these ions have an octahedral coordination geometry and therefore form structures analogous to the Cr$_7$Ni rings, but with differing ground state spins. Consequently, their potential uses may be expanded, not only to QIP data storage devices but also to magnetic refrigeration.

Usually, the paramagnetism that occurs in heterometallic wheels precludes the use of NMR spectroscopy as a characterisation tool. However, Cr$_7$Co wheels are amenable to paramagnetic NMR studies, due to the fast relaxation times of the Co which counteracts the slow relaxation times of the Cr, affording only slightly broadened peaks. Such studies may aid in the characterisation of linked-wheel systems before crystallisation, and the knowledge gained may be applied by inference to analogous Cr$_7$Ni systems.

1.5.3.3 The Carboxylate

The only restriction on the carboxylate used in the synthesis of these rings with a variety of templates is that it must be a liquid at ca. 160 °C, as a requirement of the preferred synthetic procedures. This requirement has lead to the formation of a family of wheels containing over 14 different carboxylates. If the carboxylate is not liquid at 160 °C then DMF can be used as a solvent, but the templating species will be [Me$_2$NH$_2$]$^+$. The principal carboxylate considered in this project is pivalate, as it gives the best balance between the solubility and stability of the Cr$_7$Ni wheels.

1.5.3.4 The Amine

Various amines have been used as templates for the heterometallic wheels. Altering the amine can affect the size and shape of the wheels, as the ammonium cation is always found in the centre. Cr$_7$Ni wheels have been formed around protonated primary or linear secondary amines such as di($n$-propyl)amine. It is also possible to use more complex primary or secondary amines as templates, for example using an amine such as 4-(ethylaminomethyl) pyridine can allow the synthesis of a wheel that is able to coordinate to
metal centres through its template. By moving from simple linear alkyl chains in the secondary amine to branched chains, it is possible to increase the size of the ring from octametallic to nonametallic, an example of which can be seen in Figure 12. Use of a tertiary amine has been found to lead to the formation of a decametallic wheel, of type Cr$_9$M, which can also be seen in Figure 12.

Figure 12. Examples of a nona- and deca-metallic wheel.$^{56,57}$

Cyclic metal complexes have also been utilised as templates for decametallic wheels.$^{58}$ The use of a simple aza crown species, such as 1,4,7-triazacyclononane (tacn) in a reaction otherwise identical to those carried out with other amines, allows the formation of a decametallic wheel with the complex [Ni(tacn)$_2$]$^{2+}$ as the template. Consequently, the metallic composition of these ten-membered wheels is Cr$_8$M$_2$ because the template is a dication, as confirmed by elemental analyses.

In theory, it should be possible to use these different amine templates to synthesise a full range of wheels for each combination of available trivalent and divalent metal ions.

1.5.3.5 The Fluoride

The reaction of chromium(III) nitrate with sodium pivalate in water at 40 °C allowed the formation of [Cr$_8$(OH)$_8$(O$_2$CCMe$_3$)$_{16}$] homometallic wheels.$^{59}$ The change of ligand, from fluoride to hydroxide, did not affect the structure of the wheel, with the hydroxides pointing into the centre. However, the d-d transitions of the wheel were altered
significantly and thus the hydroxide analogue is a deep purple colour instead of green. Attempts to apply this modification directly to heterometallic systems have proved unsuccessful. This is possibly due to the H-atoms of the hydroxy groups pointing into the ring centre upon formation, preventing encapsulation of the protonated amine template by H-bonding. It appears that it is this encapsulation that drives the formation of the fluoride-containing heterometallic wheels.

1.5.3.6 “Purple Wheels”

The presence of eight fluoride ions has until very recently appeared to be essential for the synthesis of these rings. It was believed that it would not be possible to form heterometallic rings of any form without fluorides bridging each metal centre. However, following the precedent with hydroxide (see above), the use of an aminopolyol as a template has led to the synthesis of a new “sugared” wheel, which can be seen in Figure 13. The polyol, either N-ethyl-D-glucamine or N-methyl-D-glucamine, becomes fully deprotonated and the resulting poly-alkoxide replaces five of the fluoride bridging ligands, producing \([\text{Cr}_7\text{NiF}_3(O_5\text{C}_8\text{H}_{14}\text{N})(O_2\text{C}^t\text{Bu})_{15}(\text{H}_2\text{O})]\) in a yield of approximately 40%. The presence of five oxygens in place of five fluorides alters the d-d transitions of the Cr\(^{III}\) ions and thus, in a similar fashion to the \([\text{Cr}_8(O\text{H})_8(\text{O}_2\text{CCMe}_3)_{16}]\) wheel, the Cr\(_7\)Ni wheels appear purple coloured.

![Figure 13. A “Sugared Wheel”, \([\text{Cr}_7\text{NiF}_3(O_5\text{C}_8\text{H}_{14}\text{N})(O_2\text{C}^t\text{Bu})_{15}(\text{H}_2\text{O})]\).](image)
The nitrogen of the “template” also coordinates to a Cr$^{III}$ ion adjacent to the divalent metal, taking up a site that should contain a bridging pivalate. Consequently a free site becomes available at the divalent metal and a solvent molecule coordinates to it. Interestingly, as the aminopolyol does not act as a template, but instead coordinates into the wheel, this wheel is neutral. This wheel has been synthesised using chromium, iron or gallium as the M$^{III}$ ion and nickel, cobalt, zinc or manganese as the M$^{II}$ ion.

The purple wheels are more anisotropic than green wheels due to their lack of symmetry. This can be observed in their Q-band EPR spectra (Figure 14). It is also possible that having a smaller number of fluorides within the structure may lead to decreased decoherence in linked wheel systems.

![Q-band, 34 GHz, 5K

Figure 14. Q-Band EPR Spectrum of a purple Cr$_7$Ni wheel.

1.5.3.7 Summary of Tuning Capabilities

When designing the wheels, it is therefore possible to use different approaches to tune not only the chemical structures of the wheels and their derivatives, but also their physical properties, as follows:

a) The ground state spin ($S$) by varying the metal ions, for examples see Table 1.
Table 1. Overall ground state spins on $\text{M}^{\text{III}}_7\text{M}^{\text{II}}$ wheels by varying metal centres.

<table>
<thead>
<tr>
<th></th>
<th>M$^{\text{II}}$</th>
<th>Ni$^{\text{II}}$</th>
<th>Co$^{\text{II}}$</th>
<th>Zn$^{\text{II}}$</th>
<th>Fe$^{\text{II}}$</th>
<th>Mn$^{\text{II}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>M$^{\text{III}}$</td>
<td>Cr$^{\text{III}}$</td>
<td>$1/2$</td>
<td>1</td>
<td>$3/2$</td>
<td>$1/2$</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>V$^{\text{III}}$</td>
<td>$0$</td>
<td>$1/2$</td>
<td>1</td>
<td>1</td>
<td>$3/2$</td>
</tr>
<tr>
<td></td>
<td>Fe$^{\text{III}}$</td>
<td>$3/2$</td>
<td>2</td>
<td>$5/2$</td>
<td>$1/2$</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Ga$^{\text{III}}$</td>
<td>$1$</td>
<td>$3/2$</td>
<td>0</td>
<td>2</td>
<td>$5/2$</td>
</tr>
</tbody>
</table>

b) The zero field splitting (D) by use of particular metal ions for example both Cr$_7$Mn and V$_7$Zn have a ground state spin of 1, but D is larger for the latter as the wheel is derived from more anisotropic ions.

c) The solubility and stability of the wheels by varying the carboxylate used as the reaction solvent. For example, using pivalic acid produces a wheel that is insoluble in acetonitrile, while using tert-butyl acetic acid produces a wheel that dissolves in acetonitrile.

1.5.4 Other Shapes in the Heterometallic Wheel Family

1.5.41 “Hourglasses”

The synthesis of heterometallic “hourglass” clusters is achieved by including a dicationic metal in the reaction, which does not adopt an octahedral coordination geometry. An example is the use of copper(II) carbonate in the presence of CrF$_3$$\cdot$4H$_2$O, pivalic acid and a secondary amine, giving the salt compound $[\text{R}_2\text{NH}_2]_2[\text{Cr}_{16}\text{Cu}_2\text{F}_{14}(\text{O}_2\text{CCMe}_3)_{22}]$ which contains a distorted dodecametallic wheel (Figure 15)."61

The copper centres are five-coordinate and connected to one Cr$^{\text{III}}$ ion through one bridging fluoride and two pivalates and to another Cr$^{\text{III}}$ ion through one fluoride and only one pivalate. The Cr$^{\text{III}}$ ion that is only doubly bridged to the Cu$^{\text{II}}$ ion also has a terminal fluoride ligand. The presence of two divalent metal ions in this distorted ring gives an overall charge of $-2$ and two protonated secondary amine cations are found in the centre of the structure, sitting in the cavities as a result of H-bonding to the centrally oriented fluoride ions.
Homometallic “horseshoes” (Figure 16) can be formed by excluding the divalent metal from the reaction. For example by reacting CrF$_3$$\cdot$4H$_2$O with a secondary amine in pivalic
acid, a pair of horseshoe-shaped clusters is formed, with an overall formula of 
\[ [R_2NH_2]_6[Cr_6F_{11}(O_2CCMe_3)]_{10} \text{(sol)} \] (sol) \[ \text{(sol)} \] \[ (R = \text{Et, } ^9\text{Pr, } ^{11}\text{C}_5\text{H}_{11}, \text{sol = solvent}). \]\(^61\) When using a bulkier secondary amine, such as iso-propyl diamine, as the horseshoe template the horseshoe clusters formed contain seven Cr\text{III} ions instead of six.

It is not yet fully understood why one template causes the formation of a hexametallic and another a heptametallic horseshoe. One possibility is that in all reactions both derivatives are formed and it is during crystallisation that the most stable arrangement for a particular template is isolated. Another possibility is that the bulk of the template has a role in the size of the horseshoe synthesised. The three linear diamines used as templates produced hexametallic horseshoes, whereas the bulkier branched diamine produced heptametallic versions, potentially due to the larger cavity size in the seven-membered horseshoe. A similar principle operates in the heterometallic wheel systems.

The crystal structures of the horseshoes showed that the arrangement of the internal Cr\text{III} ions is identical to those found in wheel structures. Each Cr\text{III} pair is bridged by two pivalate ions and one fluoride ion. The terminal Cr\text{III} sites of the horseshoe bind to the adjacent Cr\text{III} ion through two pivalate ions and one fluoride ion, but they also have three terminal fluoride ligands which are H-bonded to the protonated secondary amine cations situated between the two clusters. There are six protonated amines associated with each horseshoe pair. One amine sits in the cavity of each horseshoe and two can be found at each terminus of the horseshoe. They are held in place by H-bonding to fluorides.

1.5.4.3 “Seahorses”

A heterometallic “seahorse” cluster was formed by using a large macrocyclic templating amine, 1,4,7,10-tetraazacyclododecane (cyclen), as the template and can be seen in Figure 17.\(^58\)

Reacting cyclen with CrF\text{3}$\cdot$4H\text{2}O and nickel(II) carbonate afforded an open-chain complex with a \([\text{Ni(cyclen)}]^2^+\) unit at each end and one central Ni\text{II} ion connecting two Cr\text{6} “horseshoes” to give \([\{\text{Ni(cyclen)}\}_2\text{Cr}_{12}\text{NiF}_{20}(O_2\text{CCMe}_3)_{22}\}]\). The crystal structure showed that the twelve Cr\text{III} ions are joined together in two hexametallic chains. The coordination geometries of the internal ions of these chains are the same as in the hexametallic Cr\text{6} horseshoes. The two internal ends of the chains are linked \textit{via} two fluorides and one
pivalate bridge each to a central Ni\textsuperscript{II} ion, thus creating an “S” shape. The terminal ends of the chains are each attached to a Ni(cyclen) complex \textit{via} one fluoride and one pivalate bridge. The metal core of this molecule is virtually planar and the Ni(cyclen) complexes are held in place with a single N–H---F bond to the fluorides attached to the central Ni\textsuperscript{II} ion.

![Figure 17. A “Seahorse” Cluster Structure, $[\{\text{Ni}(\text{cyclen})\}_2\text{Cr}_12\text{NiF}_{20}(\text{O}_2\text{CCMe}_3)_22]$.\textsuperscript{58}](image)

### 1.6 LINKED WHEEL SYSTEMS

In order to fulfil the requirements needed for applications in QIP, two wheels must be linked together to form a two-qubit quantum gate. The extent of interaction between the wheels must be controllable, so studies have been undertaken to assess what factors affect this.

#### 1.6.1 Linking Wheels Indirectly through the Template

##### 1.6.1.1 Syntheses

Initial studies to link together the wheels were focussed on linking through the template. Two different methods were attempted to achieve this goal. The first method was to template the formation of two wheels around a single linker molecule, using for example a
simple linear diamine. The synthesis of this genre of linked wheel system was carried out in an identical fashion to that of a single wheel system, although the quantity of templating molecules added to the reaction was halved. Purification was achieved by recrystallisation, as the products could not be placed on a column for reasons of stability. This approach has been employed with 1,8-diaminoctane and longer chain molecules; a representative crystal structure of one of the products can be seen in Figure 18.

Figure 18. X-ray Crystal Structure of \([\{\text{H}_3\text{NC}_8\text{H}_{16}\text{NH}_3\}\{\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{16}\}_2\]^{62}.  

Another method for linking heterometallic rings involved the synthesis of a wheel around a pyridyl-functionalised amine, for example 4-(ethylaminomethyl)pyridine, followed by coordination of the pendant pyridyl units with a metal complex.\(^6\) Following the synthesis of a pyridyl-functionalised wheel, two wheels were linked together via first row transition metal centres. Preliminary studies involved the linking of wheels through copper, nickel, cobalt and ruthenium dimers\(^6\) by reaction at room temperature in toluene and in some cases diethyl ether. Gentle evaporation of the reaction solvent gave crystalline products, which in some cases could be characterised using X-ray crystallography. The crystal structure of one such assembly, \([\{\text{Cu}_2(\text{O}_2\text{CCMe}_3)_4\}\{\text{EtNH}_2\text{CH}_2\text{C}_5\text{H}_4\text{N}\}\{\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{16}\}_2\]^{62}, can be seen in Figure 19.
It was also possible to link wheels together through single metal centres, such as copper and nickel. An example of such a molecule containing copper can be seen in Figure 20.

1.6.1.2 Magnetic Studies

A working two-qubit quantum gate must display a significant and controllable electronic coupling between the magnetic components. Magnetic susceptibility and specific heat studies of all indirectly linked wheel systems have revealed negligible interaction between the two rings. Such systems therefore correspond to an “off state”, showing that the wheels can be in close proximity without affecting each other (not an unexpectedly as
there is no direct linkage between the wheels). The requirement for an “on state” has led to investigations into the formation of a direct link between two wheels.

1.6.2 Linking the Wheels Directly through the Backbone

1.6.2.1 Syntheses

Heterometallic wheels have recently been linked together directly using a novel method. For the traditional “green” wheels, a two-step process was required in order to form the links. The first step was to functionalise the wheel, by external modification of the backbone, with a linking moiety. The second step was to introduce a metal centre through which two wheels may be joined together using coordination chemistry.

The general synthetic procedure for such a substitution involves heating the wheel with an approximately four-fold excess of the incoming carboxylate in propan-1-ol under reflux. The carboxylic acids are not very soluble in propan-1-ol, so the reaction takes place over approximately 24 h. The first substitution reaction was attempted with nicotinic acid as it has a relatively good solubility. However, isonicotinic acid (ISNA) substitutes into the wheel more efficiently, so this project focuses on ISNA-functionalised wheels.

This substitution process was selective, as the Ni^{II} ion is much more labile than the relatively inert Cr^{III} ions, so both the primary and secondary substitutions occur at the nickel and one of the adjacent chromium ions. Column chromatography on silica gel was used to isolate the product, eluting initially with toluene to bring off the unsubstituted wheel, then increasing the gradient using a 40:3 mixture of 40/60 °C petroleum ether and ethyl acetate to bring off products with higher degrees of substitution. With ISNA, both the mono and disubstituted products have been isolated, characterised and used for further synthetic studies. Crystal structures of these substituted wheels can be seen in Figure 21.
Microwave and solvothermal techniques have also been used to achieve substitution of carboxylates into the wheels, but these approaches caused more decomposition than standard methods. Different solvents can be used for reactions of wheels with other carboxylate groups; for example, using tert-butyl acetate in place of pivalate groups allows the substitution reactions to be carried out in acetonitrile.

By altering the pendant group on the incoming carboxylate, it is possible to alter the chemistry that can be achieved with the substituted product. Using a pyridyl unit allows the wheel to act as a potential bulky ligand, while using an alkyne substituent allows the wheel to undergo ‘Click chemistry’ to join together two wheels. Thiol substituents may allow the wheels to attach directly to gold surfaces in self-assembled monolayer formations.
The ISNA-substituted wheels can act as bulky ligands with first row transition metal centres.\(^6^5\) Reaction of such a wheel with \([\text{Cu}_2(\text{O}_2\text{CCMe}_3)_4(\text{HO}_2\text{CCMe}_3)_2]\) at room temperature in toluene afforded a linked wheel system (Figure 22). Heating the ISNA-substituted wheels with \(\text{Cu(NO}_3\text{)}_2\cdot2.5\text{H}_2\text{O}\) in acetone allowed the formation of another type of linked wheel system (Figure 23).

![Figure 23. Wheels linked directly through a single copper centre.\(^6^5\)](image)

Reaction of a disubstituted wheel with a metal complex, such as \([\text{Cu}_2(\text{O}_2\text{CCMe}_3)_4(\text{HO}_2\text{CCMe}_3)_2]\), allows the formation of a polymeric structure where two metal complexes are linked together through a wheel.\(^6^6\) An example of such a structure can be seen in Figure 24.

![Figure 24. Polymer formed by disubstituted wheels linked through copper dimers.\(^6^6\)](image)
1.6.2.2 Magnetic Studies

EPR studies on a system with two ISNA-substituted wheels joined through a copper dimer showed no interaction between the wheels, because the spectrum is identical to that of the uncomplexed wheel (Figure 25).\textsuperscript{65} This result is predictable as the two paramagnetic copper centres in the dimer are very strongly anti-ferromagnetically coupled, forming a diamagnetic, partially conjugated moiety that prevents any communication between the two wheels.

Figure 25. Q-Band EPR Spectra of Individual Cr\textsubscript{7}Ni Ring (black) and 2 Cr\textsubscript{7}Ni wheels linked through a Cu dimer (blue).\textsuperscript{65}

When the Cr\textsubscript{7}Ni wheels are linked through a single copper centre, however, a very different EPR spectrum was observed.\textsuperscript{65} The spectrum of the linked wheel system, shown in red in Figure 26, was much more complex than the simple singlet observed for a single Cr\textsubscript{7}Ni wheel. This observation is due to the paramagnetic nature of the Cu centre and the conjugated links between the two Cr\textsubscript{7}Ni wheels allowing interactions between them.
It has therefore been demonstrated that by linking the wheels together directly it is possible to observe both the “on” and “off” states required for uses in QIP. However, for such an application, the “on” and “off” states must be generated within the same molecule.

1.6.3 Linking Purple Wheels together

While it is plausible that purple wheels could be linked together directly in a manner identical to that used for green wheels, a novel route was considered taking advantage of the spare coordination site available on the divalent metal centre. Molecules containing two or more pendant pyridyl groups have been used to link wheels together, with the pyridyl coordinating to the divalent metal.

1.6.3.1 Syntheses

Reactions of a purple cluster with 4-picoline (4-pic) in acetone allowed substitution of the coordinated water molecule to produce \([\text{Cr}_7\text{NiF}_3(\text{O}_2\text{C}_8\text{H}_{14}\text{N})(\text{O}_2\text{C}^\text{tBu})_5(4\text{-pic})]\) which has also been structurally characterised, confirming the coordination of 4-pic to the \(\text{Ni}^{II}\) centre. This concept was then extended to molecules containing two pendent pyridyl groups, such as 4,4′-bipyridine (bpy) and \(\text{trans}-1,2\)-bis(4-pyridyl)ethylene (bpe), in order
to form linked wheel systems. The synthetic procedures used are straightforward and involve a short period of heating in a low boiling point solvent such as acetone or dichloromethane. Once the linked wheel systems were formed they were isolated by crystallisation, and characterised by X-ray crystallography and elemental analyses.

1.6.3.2 Magnetic Studies

Systems containing wheels linked through organic moieties did not show any difference in their SQUID measurements. This is to be expected as the interactions between the two wheels are relatively weak, and more significant interactions are required to affect SQUID measurements. EPR spectroscopy can detect much weaker interactions and consequently a difference in spectra may be observed between two wheels linked through bpy and an individual wheel (Figure 27). The spectrum of the linked wheel system, corresponds to an S = 1 ground state below 10 K, meaning the wheels are coupled ferromagnetically to each other.

![EPR spectrum of unlinked purple Cr7Ni wheels (black) and linked through bpy (red).](image)

**Figure 27.** EPR spectrum of unlinked purple Cr7Ni wheels (black) and linked through bpy (red).\(^{60}\)

1.7 ATTACHING WHEELS TO SURFACES

In order to apply these systems, it will be necessary to attach them to surfaces. Consequently, studies have been carried out to ascertain the suitability of different methods for this purpose. Further to investigations with Mn\(_{12}\) clusters,\(^ {67}\) the viability of attaching Cr7Ni wheels to gold surfaces has been studied.\(^ {68,69}\) Wheels were deposited on Au(111)
surfaces in a one-step procedure, by immersing a 5 mM solution in either tetrahydrofuran or dichloromethane over 10 minutes. The treated surfaces were then studied using scanning tunnelling microscopy (STM) and X-ray photoemission spectroscopy (XPS).

Figure 28. Thio-functionalised wheels for attachment to gold surfaces; (a) and (b) thiols in template, (c) and (d) thiols in backbone.

Four thio-functionalised wheels (Figure 28) and three other functionalised wheels (Figure 29) have been used for these experiments; three of these (a, b and e) contained the surface-binding functionality in the template of the wheel and four were functionalised in their backbones (c, d, f and g). The former showed inefficient grafting, with only 2–10% coverage of the surface area by wheels; this was primarily caused by the wheels detaching from their templates, leaving a self-assembled monolayer (SAM) of the templates alone. However, the backbone-functionalised wheels adsorbed much more successfully, giving up to 35% surface coverage.

In keeping with the known high affinity of sulfur for gold, the thio-functionalised wheels adsorbed most strongly, but reasonable surface coverages were nevertheless observed for the other wheels. Physical studies showed that the electronic properties of the surface-
attached wheels remained largely unchanged when compared with their behaviour in solution. However, surface binding did cause a reduction in the coupling constants between Cr and Ni, which in turn lowered the energy gap between the $S = \frac{1}{2}$ ground state and other excited states.

![Figure 29](image1.png)

Figure 29. Non-thio functionalised wheels for gold surface studies.

Wheels have also been adsorbed onto highly-oriented pyrolytic graphite (HOPG). The wheels synthesised for this purpose contained large alkyl or benzyl groups (Figure 30), which can bind to HOPG by van der Waal’s (VdW) or π-π interactions. Two techniques were used in this study, the first being a one-step method whereby the substrate is dipped into a toluene solution of the wheel. The wheels adsorbed readily onto the HOPG surface on bulk, but using $10^{-4}$ M concentrations gave only sub-monolayers. The second technique involved two-steps. First, a SAM of the salt CH$_3$(CH$_2$)$_{15}$SO$_3$Na was applied to the surface, then a cationic wheel was bound by interaction with the anionic $-\text{SO}_3^-$ groups. The electrostatic bonds formed were strong and the wheels remained on the surface, even after a strong rinse with dichloromethane.
1.8 FORMING NOVEL LINKED WHEEL SYSTEMS

1.8.1 Non-Switchable Systems

While it is desirable to link wheels together through molecules that are able to act as a switch for quantum gate applications, it is also important to be able to understand how wheels interact when linked through various moieties. Therefore, some target linker molecules were identified, primarily for studying the interactions they mediate as opposed to creating switchable systems.

In order for qubits to become entangled, there must be a direct path for magnetic interactions, so one line of investigation considered is fully conjugated organic molecules. As purple wheels have already been linked together and shown to interact through 4,4’-bpy and bpe, extended versions of these molecules are viable options for probing how the length of the linkage affects the communication between the wheels (Figure 31).
In the case of green wheels linked directly by using an ISNA-functionalised carboxylate in the backbone, it has been possible to observe interaction by linking through a paramagnetic metal centre. Insertion of a diamagnetic centre between the two wheels acts as a break in the circuit and prevents any interaction. There are two variables by which we can further study these wheels. The first is to attempt substitution reactions with other functionalised carboxylates (Figure 32) that may be able to act as ligands to metal centres. By comparing EPR spectra of analogous ISNA compounds, it may be possible to probe the effects of varying the carboxylate on the entanglement of the wheels. The second variable is the transition metal centre. So far, wheels have been linked together directly using copper and copper dimers. It would be interesting to observe what effect other metal centres will have on the interaction occurring between the wheels. Some of the metal centres considered for linking wheels are; $\text{M}^{II}(\text{CF}_3\text{COCHCOCF}_3)_2$ where $\text{M}^{II} = \text{Cu}, \text{Ni}$ and $\text{Mn}$, fac-ReCl(\text{CO})$_3$ and the triangular system, $\text{Fe}_2\text{NiO(O}_2\text{CCMe}_3)$_6$.

The final area of consideration for non-switchable systems is that of mixed-wheel systems. It has been shown that green wheels with pyridyl functionalised carboxylates may act as ligands to transition metal centres.$^{65}$ The purple wheel has a free coordination site on the divalent metal centre, so it may be possible to attach a green wheel to a purple wheel simply by adding them together.
1.8.2 Switchable Systems

Many recent studies have focused on the concept of “molecular switches”, molecules that can exist in two or more stable states. Conversion between these states can be achieved by various means; either by exploiting photoexcitation\textsuperscript{72,73} or redox activity,\textsuperscript{74} or by alteration of other external conditions, such as pH\textsuperscript{75} or temperature.\textsuperscript{76} An example of a photoswitchable compound can be seen in Figure 33. The orange bead-like molecule can sit around either the green or the purple functional group dependent on its environment.\textsuperscript{77}

![Figure 33. An example of a photoswitchable rotaxane synthesised by Stoddart et al.\textsuperscript{77}](image)

A relevant example of a molecular switch was published in 2000 by Irie and Matsuda, who showed that it is possible to control the interaction between two radicals by linking them through a diarylethene molecule.\textsuperscript{78} Exposure of the sample to UV light, caused a ring closing reaction to occur, allowing full conjugation in the linking molecule (Figure 34). Differences were observed in the magnetic SQUID measurements and electrochemistry of the two compounds as a result of intramolecular anti-ferromagnetic interactions.

![Figure 34. Irie’s molecular switch involving an intramolecular magnetic interaction.\textsuperscript{78}](image)

This project is particularly focused on the use of photo-excitable or redox-active compounds as switches that may be able to modulate interaction between the wheels. Systems to be considered include viologens, [M(bpy)\textsubscript{3}]\textsuperscript{2+} (M = Fe or Ru) and fac-ReCl(CO)\textsubscript{3}(bpy)-based MLCT chromophores.
1.8.2.1 Viologens

The viologens are a series of bis-quaternised derivatives of 4,4'-bpy, which can undergo two reversible one electron reductions (Figure 35). The first reduction produces a monoradical cation, which is an intense blue colour,\textsuperscript{79} and further reduction produces a quinoid. Synthesis is generally achieved by refluxing a halogenated alkane (RX) with 4,4'-bpy in a polar solvent to allow the bis-quaternisation to occur. Use of a less polar solvent allows the formation of the mono-quarternised product. Viologens have proved to be useful in many areas, such as solar energy conversion, herbicides, electrochromic devices and molecular electronics,\textsuperscript{80} and have been tested recently as molecular switches for quantum computing purposes, by controlling electron transport between gold nano-particles and an electrode surface.\textsuperscript{81} This type of molecule (Figure 36) will only be considered for linking wheels indirectly due to the synthetic limitations inherent in attaching cationic species to the backbone of wheels.

1.8.2.2 Photo- and Redox-Active Metal Complexes

Another type of switchable system considered for use as a linking molecule between the wheels is a MLCT-active complex. In such complexes, charge-transfer can occur upon excitation in the UV or short wavelength visible light regions allowing an electron to be promoted from a d-orbital on the metal centre to a low-lying $\pi^*$ anti-bonding orbital on the ligand. These transitions are both Laporte and spin allowed, so their molar extinction coefficients are in the region of $10^4$, and the majority of MLCT chromophores are very strongly coloured.
There are many types of MLCT chromophore. However, this project will focus on ruthenium polypyridyl and rhenium α-diimine systems (Figures 37 and 38) as these are very well studied, and known to have long excited-state lifetimes that may be sufficient for our purposes. In both cases the diamagnetic ground-state, low spin $d^6$ metal centres ($\text{Ru}^{II}$ or $\text{Re}^I$) lose an electron temporarily upon MLCT excitation and become paramagnetic with a low spin $d^5$, $S = \frac{1}{2}$, configuration. This change from diamagnetic to paramagnetism should greatly affect any interactions between the wheels.

The photochemical properties of $[\text{Ru}(\text{bpy})_3]^{2+}$ have been very well studied. The UV-vis spectrum of this complex has two absorptions due to ligand-based excitations at 185 and...
285 nm, and two MLCT transitions at 240 and 450 nm. If excited at these wavelengths, long-lived luminescence (on the 100–3000 ns scale) is observed. Another much used MLCT chromophore is fac-ReCl(CO)₃(bpy), which shows an MLCT absorption band at 370 nm. The lifetime of this excited state (on the 10–500 ns scale) is shorter than that of [Ru(bpy)₃]²⁺ because of intramolecular quenching, caused by the chloride ligand. The advantage of both of these types of complexes is that the α-diimine ligands can be altered readily, allowing control over the absorption and emission properties.

The ruthenium complexes synthesised should also show a reversible Ru²⁺/³⁺ oxidation which can be observed by electrochemical techniques. Replacement of the ruthenium centre with iron will also give reversible redox properties, although Fe²⁺ has lower $E_{1/2}$ values and is thus more readily oxidised than Ru²⁺. Chemical or electrochemical redox is clearly related to MLCT and may provide an alternative method of switching the interaction between the wheels.

**1.9 ANALYTICAL TECHNIQUES**

**1.9.1 Paramagnetic ¹H NMR Spectroscopy**

In general, NMR spectroscopy is an inappropriate analytical technique for systems containing Cr₇M wheels as the overly fast relaxation of the paramagnetic Cr³⁺ ions broadens the spectrum severely. However, it is possible to use NMR spectroscopy with the Cr₇Co analogues of the wheels, as the slow relaxation of the Co²⁺ ion counteracts the effects of the Cr³⁺ ions and thus a spectrum may be observed. Previous studies have been carried out on simple symmetrical Cr₇Co wheels, demonstrating that a degree of information can be obtained.
The spectra obtained are a lot broader than expected for a regular diamagnetic sample, and consequently splitting of the peaks cannot be observed. The chemical shifts of the peaks, instead of being restricted between 0 and 10 ppm, are changed greatly and can be found between –80 and 15 ppm. Also, the integration of the peaks is not always very accurate. Although this technique is not ideal, it can prove useful to probe how a paramagnetic molecule is arranged in solution, while X-ray crystallography is applicable to the solid state only.

1.9.2 EPR Spectroscopy

EPR spectroscopy is a technique used for studying paramagnetic species. It is very similar to NMR spectroscopy, but probes electron- as opposed to nuclear-spin transitions. EPR can be studied by varying either the magnetic field or the frequency of radiation, but in general the frequency is kept the same while the magnetic field varies. At The University of Manchester, a number of machines of different frequency are used to run EPR spectra; X-band (8–10 GHz), K-band (24 GHz), Q-band (35 GHz) and W-band (95 GHz).

Samples can be run in a variety of states; gases, single crystals, solutions, powders and frozen solutions. The spectrum is usually shown as a first derivative of the absorption spectrum, to reduce noise seen on the absorption spectrum.

The g-factor, which is a unitless measurement of the magnetic moment of a particular electron (the moment of a free electron is 2.0023), and analogous to the chemical shift in NMR spectrum can be calculated using Equation 6, where \( h \) is Planck’s constant \( (6.626 \times 10^{-34} \text{ J s}^{-1}) \), \( \nu \) is the frequency of radiation \( (\text{s}^{-1}) \), \( \beta \) is the Bohr magnetron \( (9.274 \times 10^{-24} \text{ J T}^{-1}) \) and \( B \) is the strength of the magnetic field \( (\text{T}) \).

\[
\text{Equation 6:} \quad g = \frac{h \nu}{\beta B}
\]

Hyperfine splitting is possible in EPR spectra, either through coupling of the electron to its own nucleus’ spin or through coupling to that of a nearby nucleus. As in NMR spectra, the coupling ratio follows Pascal’s triangle of intensities. The hyperfine constant, \( a \), which is the distance between two of the peaks can be converted into a value, \( A \), in Hz by using Equation 7.
Another factor that can affect the EPR spectrum of a molecule is anisotropy, i.e. orientation-dependence. The g-factor is split into three components: $g_x$, $g_y$ and $g_z$. If $g_x = g_y = g_z$, the spectrum is considered to be isotropic. When $g_x = g_y \neq g_z$, the spectrum is said to be axial. If $g_z$ is larger than $g_x$ and $g_y$, then a small peak is seen at low field and a large peak is seen at high field. If $g_z$ is smaller than $g_x$ and $g_y$, a large peak is observed at low field and a small peak at high field. When $g_x \neq g_y \neq g_z$, the spectrum is said to be rhombic and 3 large peaks are observed, corresponding to the different values of g. The wheels under study here are axially symmetrical and $g_z$ is smaller than $g_{xy}$.

1.10 INITIAL AIMS OF THE PROJECT

The aims of this project are to extend the series of linked wheel systems in order to better understand how the link affects interaction between the wheels. This is to be investigated by three different synthetic strategies.

1. Functionalisation of the wheel followed by coordination to a metal centre.

Functionalisation of green wheels may be achieved by selective substitution of a pivalate group for a carboxylate containing a pendant coordinating group. The product may then be used as a bulky ligand in the synthesis of transition metal complexes, forming linked wheel systems. A number of carboxylates will be used to assess their suitability as substituents and their behaviour as bulky ligands in previously used metal complexes. The series of metal complexes used with these wheels will also be expanded to include some potential switchable MLCT active Ru$^{II}$ and Re$I$ complexes.

2. Coordination of pyridyl to a vacant site on the divalent metal in purple wheels.

As previously discussed in Section 1.6.3, linking of purple wheels is carried out by coordinating a pyridyl ligand to the spare site on the divalent metal. Attempts will be made to link wheels using more lengthy organic molecules with the aim of studying how the interaction between the wheels is affected by the length of the link. Another avenue of
investigation to be considered is the linking of purple wheels through some MLCT active
transition metal complexes based upon quaterpyridyl ligands.

3. Linking the wheels through hydrogen-bonds to a template molecule.

Wheels can be linked indirectly either through direct formation of wheels about a pre-made
template containing two amine groups or by using a wheel templated about a pyridyl
containing amine as a ligand in a coordination complex. A variety of switchable
templating molecules will be synthesised in order to attempt direct synthesis of linked
wheel systems.
Chapter 2 – Linked Green Wheel Systems
2.1 INTRODUCTION

The attempted formation of two qubit quantum gates using Cr\textsubscript{7}Ni heterometallic wheels first began by linking the wheels together through their templates.\textsuperscript{62} While some success was achieved in the synthesis of these systems, the properties desired for use in quantum computing were not observed.\textsuperscript{64} Consequently an alternative method was devised, whereby the wheels were linked together directly by replacing one of the pivalates with a carboxylate with a pendent coordinating group such as a pyridyl ring. To date, the isonicotinate-substituted wheel \([\text{Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPy})]\), \textsuperscript{203}, has been the most used as it can be produced in a reasonable yield and has been proven to act as a bulky ligand in simple coordination chemistry under mild conditions.\textsuperscript{65}

Magnetic studies on two different copper complexes demonstrated that by linking the wheels together directly it is possible to obtain the two states (ON/OFF) required in a two-qubit quantum gate.\textsuperscript{65} The first complex, a copper dimer \(\text{Cu}_2(\text{O}_2\text{CCMe}_3)_4(\text{203})_2\), shows no interaction of spins of the wheels. However, the second copper complex, \(\text{Cu(NO}_3)_2(\text{H}_2\text{O})_2(\text{203})_2\), shows a coupling of the wheels, observable in the EPR spectrum (Figure 39).

![EPR spectrum of individual wheel (black) versus Cu linked-wheel system (red).\textsuperscript{65}](image-url)
The work in this chapter includes the further development of the series of mono- and disubstituted wheels, and the resultant metal complexes that can be obtained by using them as bulky ligands. Studies have been undertaken into the abilities of 203 as a ligand under more harsh conditions, for use with second and third row transition metal centres. Of particular interest in this respect are Ru$^{II}$ and Re$^{I}$ complexes, which are well known to demonstrate MLCT excited states with relatively long lifetimes, that may enable an ON/OFF state switching to occur.
2.2 EXPERIMENTAL PROCEDURES

2.2.1 Materials and Procedures

The compounds 
\[ \text{[Co(O}_2\text{CCMe}_3)_2]n \]
\[ \text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_1]_5 \]
\[ \text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{CoF}_8(\text{O}_2\text{CCMe}_3)_1]_5 \]
\[ \text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_1]_5(\text{O}_2\text{CPy}) \]
\[ \text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_1]_5(\text{O}_2\text{CPy})(\text{HO}_2\text{CCMe}_3)_3 \]
\[ \text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_1]_5(\text{O}_2\text{CPy})(\text{HO}_2\text{CCMe}_3)_3 \]

were synthesised by literature methods. All other materials were purchased commercially and used as supplied.

2.2.2 Physical Measurements

All studies were carried out at the University of Manchester unless stated otherwise. Paramagnetic 1H NMR spectra were run by Dr Louise Natrajan and Mr Thomas Faust on a Bruker Advance 400 MHz spectrometer. IR data were recorded using a Varian Bio-Rad Excalibur series IR instrument and UV data were recorded on a Shimadzu UV-2401 PC UV-VIS spectrophotometer. Elemental analyses were performed by the Microanalytical Laboratory, and mass spectra were obtained using electrospray analysis on a Micromass Platform spectrometer. Linked wheel systems were sent to Dr Lindsay Harding at the University of Huddersfield for analysis using a Bruker MicroTOF-Q mass spectrometer with an electrospray ionisation source. EPR studies were performed by Dr Floriana Tuna and Mr John Machin; Q-Band EPR was run on an Elecsys E500 spectrometer equipped with a Bruker 2 tesla magnet, and variable temperature studies used an Oxford VT liquid helium cryostat between 5 and 30 K.

2.2.3 Syntheses

\[ \text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_1]_5(\text{O}_2\text{CCCHCH-4-Py}) \]

\[ \text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_1]_6 \]

(5.035 g, 2.19 mmol) and trans-3-(pyrid-4-yl)acrylic acid (4-pyac) (1.291 g, 8.77 mmol) were refluxed in propan-1-ol (100 ml) for 19.5 h before the solvent was removed under vacuum. The impure product was extracted into diethyl ether,
the solution filtered and the solvent removed under vacuum. The solid was then washed with acetonitrile and dried thoroughly before further purification by column chromatography on silica column, initially eluting with toluene to remove the starting material. The eluent was then changed to a 20:1 mixture of 40–60 °C petroleum ether/ethyl acetate to separate the mono-substituted wheel from more highly substituted products. Once a band had become fully distinguishable, a gradient up to 8:1 40–60 °C petroleum ether/ethyl acetate was used to remove the desired product as a green solid. Yield: 0.628 g (12%).

+ve electrospray: \( m/z: 2342 \ [MH]^+, 2364 \ [MNa]^+ \).  
-ve electrospray: \( m/z: 2340 \ [M – Pr_2NH_2]^-, 2239 \ [M – Pr_2NH_2]^-. \)  
Elemental analysis calcd. (%) for C_{89}H_{157}Cr_{7}F_{8}N_{2}NiO_{32}: C 45.65, H 6.76, N 1.20, Cr 15.54. Found: C 45.40, H 6.88, N 1.07, Cr 15.49.

\[ \text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CCHCH-3-Py})] \ 205 \]

This compound was prepared and purified in manner similar to 204 by using [Pr2NH2][Cr7NiF8(O2CCMe3)16] (5.093 g, 2.22 mmol) and trans-3-(pyrid-3-yl)acrylic acid (1.288 g, 8.75 mmol) with a reflux time of 20 h, giving a green solid. Yield: 0.335 g (6%).

+ve electrospray: \( m/z: 2342 \ [MH]^+, 2364 \ [MNa]^+ \).  
-ve electrospray: \( m/z: 2340 \ [M – \text{Pr}_2\text{NH}_2]^-, 2340 \ [M – \text{Pr}_2\text{NH}_2]^-. \)  
Elemental analysis calcd. (%) for C_{89}H_{157}Cr_{7}F_{8}N_{2}NiO_{32}: C 45.65, H 6.76, N 1.20, Cr 15.54. Found: C 45.80, H 6.92, N 1.09, Cr 15.18.

\[ \text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPyd})] :\text{Pyd} = \text{4-pyridazine} \ 206 \]

This compound was prepared and purified in manner similar to 205 by using [Pr2NH2][Cr7NiF8(O2CCMe3)16] (5.10 g, 2.22 mmol) and 4-pyridazine carboxylic acid (4-pyd) (1.063 g, 8.57 mmol), giving a green solid. Yield: 2.27 g (44%).

+ve electrospray: \( m/z: 2316 \ [MH]^+, 2339 \ [MNa]^+ \).  
-ve electrospray: \( m/z: 2214 \ [M – \text{Pr}_2\text{NH}_2]^-, 2214 \ [M – \text{Pr}_2\text{NH}_2]^-. \)  
Elemental analysis calcd. (%) for C_{86}H_{134}Cr_{7}F_{8}N_{3}NiO_{32}: C 44.58, H 6.70, N 1.81, Cr 15.71. Found: C 44.70, H 7.05, N 1.76, Cr 15.52.

\[ \text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{CoF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPy})] \ 207 \]

This compound was prepared and purified in manner similar to 205 by using [Pr2NH2][Cr7CoF8(O2CCMe3)16] (5.118 g, 2.23 mmol) and ISNA (0.836 g, 6.79 mmol) with a reflux time of 17 h, giving a green solid. Yield: 1.24 g (24%).
2316 [MH]$^+$, 2339 [MNa]$^-$. Elemental analysis calcd. (%) for C$_{87}$H$_{155}$CoCr$_7$F$_8$N$_2$O$_3$: C 45.12, H 6.75, N 1.21, Cr 15.72. Found: C 44.65, H 6.73, N 1.17, Cr 16.11.

$[\text{Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{14}(\text{O}_2\text{CPy})_2]$ 209

This compound was prepared and purified in manner similar to 206, but the desired product was eluted from the column after the mono-substituted product by using 20:3 40–60 °C petroleum ether/ethyl acetate, giving a green solid. Yield: 310 mg (6%). +ve electrospray: $m/z$: 2361 [MNa]$^+$. –ve electrospray: $m/z$: 2335 [M – Pr$_2$NH$_2$], 2338 [M – H$^-$]. Elemental analysis calcd. (%) for C$_{86}$H$_{148}$CoCr$_7$F$_8$N$_5$O$_3$: C 45.43, H 6.60, N 2.82, Cr 14.66. Found: C 45.88, H 6.90, N 2.61, Cr 14.23.

$[\text{Pr}_2\text{NH}_2][\text{Cr}_7\text{CoF}_8(\text{O}_2\text{CCMe}_3)_{14}(\text{O}_2\text{CPy})_2]$ 210

This compound was prepared and purified in a manner similar to 207, but the desired product was eluted from the column after the mono-substituted product by using 20:3 40–60 °C petroleum ether/ethyl acetate, giving a green solid. Yield: 143 mg (3%). +ve electrospray: $m/z$: 2359.6 [MNa]$^+$. Elemental analysis calcd. (%) for C$_{88}$H$_{150}$CoCr$_7$F$_8$N$_2$O$_7$: C 45.23, H 6.47, N 1.80, Cr 15.57. Found: C 45.36, H 6.69, N 1.65, Cr 15.24.

Cu$_2$(O$_2$CCMe$_3$)$_4$([Pr$_2$NH$_2$][Cr$_7$NiF$_8$(O$_2$CCMe$_3$)$_{15}$(O$_2$CCHCH-4-Py)])$_2$ 211

Cu$_2$(O$_2$CCMe$_3$)$_4$(HO$_2$CCMe$_3$)$_2$ (38.0 mg, 0.052 mmol) was added to a solution of 204 (253 mg, 0.108 mmol) in toluene (10 ml) and the mixture was stirred at room temperature for 15 min. The mixture was refluxed for 5 min, cooled to room temperature and stirred for 95 min then covered and left to stand overnight. Slow evaporation of the solvent at room temperature afforded the green crystalline product. Yield: 38.8 mg (14%). Elemental analysis calcd. (%) for C$_{198}$H$_{350}$Cr$_{14}$Cu$_2$F$_{16}$N$_8$Ni$_2$O$_{72}$: C 45.60, H 6.76, N 1.07, Cr 13.96, Cu 2.44. Found: C 45.70, H 6.85, N 1.02, Cr 13.59, Cu 2.48.

Cu$_2$(O$_2$CCMe$_3$)$_4$([Pr$_2$NH$_2$][Cr$_7$NiF$_8$(O$_2$CCMe$_3$)$_{15}$(O$_2$CCHCH-3-Py)])$_2$ 212

This compound was prepared and purified in manner similar to 211 by using Cu$_2$(O$_2$CCMe$_3$)$_4$(HO$_2$CCMe$_3$)$_2$ (38.6 mg, 0.052 mmol) and 205 (252 mg, 0.108 mmol),
Chapter 2 – Linked Green Wheel Systems

giving a green crystalline product. Yield: 78 mg (64%). Elemental analysis calcd. (%) for C_{198}H_{350}Cr_{14}Cu_{2}F_{16}N_{4}Ni_{2}O_{72}: C 45.60, H 6.76, N 1.07, Cr 13.96, Cu 2.44. Found: C 43.76, H 6.81, N 0.92, Cr 13.93, Cu 2.37.

\[
\text{Cu}_2(\text{O}_2\text{CCMe}_{15})_4(\text{[Pr}_2\text{NH}_2])\text{[Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPyd})])_2
\]

This compound was prepared and purified in manner similar to 211 by using Cu_{2}(O_2CCMe)_4(HO_2CCMe)_2 (38.8 mg, 0.053 mmol) and 206 (250 mg, 0.108 mmol), giving a green crystalline product. Yield: 125 mg (44%). Elemental analysis calcd. (%) for C_{192}H_{344}Cr_{14}Cu_{2}F_{16}N_{6}Ni_{2}O_{72}: C 44.65, H 6.71, N 1.63, Cr 14.09, Cu 2.46. Found: C 44.78, H 6.84, N 1.51, Cr 13.27, Cu 2.47.

\[
\text{Cu(NO}_3)_2(\text{H}_2\text{O})(\text{[Pr}_2\text{NH}_2])\text{[Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CCHCH-4-Py})])_2
\]

Cu(NO_3)_2•2.5H_2O (11.6 mg, 0.050 mmol) was added to a warm solution of 204 (302 mg, 0.129 mmol) in acetone (40 ml). The reaction was stirred for 5 min then filtered. The flask and sinter were washed with further acetone (5 ml) and the solution left covered at room temperature for 48 h, affording green crystals. Yield: 126 mg (52%). Elemental analysis calcd. (%) for C_{178}H_{316}Cr_{14}CuF_{28}N_{4}Ni_{2}O_{71}: C 43.73, H 6.51, N 1.72, Cr 14.89, Cu 1.30, Ni 2.40. Found: C 43.51, H 6.55, N 1.71, Cr 14.94, Cu 1.32, Ni 2.35.

\[
\text{Cu(hfac)}_2(\text{[Pr}_2\text{NH}_2])\text{[Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPy})])_2
\]

To a solution of 203 (101 mg, 0.043 mmol) in refluxing acetone (20 ml) was added Cu(hfac)_2(H_2O)_2 (10.8 mg, 0.021 mmol). The solution was heated with stirring for 5 mins during which time the product formed as a green precipitate, which was collected by filtration, washed with acetone and dried in air. Yield: 46 mg (43%). Elemental analysis calcd. (%) for C_{184}H_{312}Cr_{14}CuF_{28}N_{4}Ni_{2}O_{68}: C 43.25, H 6.15, N 1.10, Cr 14.25, Cu 1.24, Ni 2.30. Found: C 43.15, H 6.18, N 1.11, Cr 13.88, Cu 1.23, Ni 2.29.

\[
\text{Ni(hfac)}_2(\text{[Pr}_2\text{NH}_2])\text{[Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPyd})])_2
\]

To a solution of 206 (103 mg, 0.044 mmol) in warm acetone (15 ml) was added a solution of Ni(hfac)_2(H_2O)_2 (10.8 mg, 0.021 mmol) in acetone (5 ml). The reaction was heated for 5 min then cooled to room temperature, covered and left for 24 h. The resulting green
crystals were collected by filtration, washed with a small amount of acetone and dried under vacuum. Yield: 42 mg (39%). Elemental analysis calcd. (%) for C\textsubscript{182}H\textsubscript{310}Cr\textsubscript{14}F\textsubscript{28}N\textsubscript{6}Ni\textsubscript{3}O\textsubscript{68}: C 42.81, H 6.12, N 1.65, Cr 14.26, Ni 3.45. Found: C 42.47, H 6.00, N 1.59, Cr 13.85, Ni 3.46.

\textit{Mn(hfac)}\textsubscript{2}([Pr\textsubscript{2}NH\textsubscript{2}][Cr\textsubscript{7}NiF\textsubscript{8}(O\textsubscript{2}CCMe\textsubscript{3})\textsubscript{15}(O\textsubscript{2}CPyd)])\textsubscript{2} 218

This compound was prepared and purified in manner similar to 218 by using 206 (106 mg, 0.046 mmol) and Mn(hfac)\textsubscript{2}(H\textsubscript{2}O)\textsubscript{2} (14.4 mg, 0.031 mmol) in place of Ni(hfac)\textsubscript{2}(H\textsubscript{2}O)\textsubscript{2}. The crystalline product was collected after 48 h at room temperature. Yield: 56 mg (47%). Elemental analysis calcd. (%) for C\textsubscript{182}H\textsubscript{310}Cr\textsubscript{14}F\textsubscript{28}Ni\textsubscript{2}O\textsubscript{68}•2Me\textsubscript{2}CO: C 43.27, H 6.22, N 1.61, Cr 13.95, Mn 1.05. Found: C 43.15, H 6.19, N 1.66, Cr 13.64, Mn 0.80.

\textit{Fe\textsubscript{2}NiO(O\textsubscript{2}CCMe\textsubscript{3})\textsubscript{6}([Pr\textsubscript{2}NH\textsubscript{2}][Cr\textsubscript{7}NiF\textsubscript{8}(O\textsubscript{2}CCMe\textsubscript{3})\textsubscript{15}(O\textsubscript{2}CCHCHPy)])\textsubscript{3} 219

Fe\textsubscript{2}NiO(O\textsubscript{2}CCMe\textsubscript{3})\textsubscript{6}(HO\textsubscript{2}CCMe\textsubscript{3})\textsubscript{3} (26.4 mg, 0.024 mmol) and 204 (101 mg, 0.043 mmol) were dissolved in n-hexane (10 ml) and stirred at room temperature for 20 h before the solvent was allowed to evaporate to dryness. Crystals suitable for X-ray crystallography were obtained by evaporation of a diethyl ether/acetonel solution at room temperature. Yield: 71 mg (63 %). Elemental analysis calcd. (%) for C\textsubscript{297}H\textsubscript{525}Cr\textsubscript{21}F\textsubscript{28}Fe\textsubscript{2}N\textsubscript{4}Ni\textsubscript{4}O\textsubscript{109}: C 45.62, H 6.77, N 1.07, Cr 13.97, Ni 3.00. Found: C 43.76, H 6.65, N 0.93, Cr 12.26, Ni 3.02.

\textit{ReCl(CO)}\textsubscript{3}([Pr\textsubscript{2}NH\textsubscript{2}][Cr\textsubscript{7}NiF\textsubscript{8}(O\textsubscript{2}CCMe\textsubscript{3})\textsubscript{15}(O\textsubscript{2}CPy)])\textsubscript{2} 220

ReCl(CO)\textsubscript{3} (30.9 mg, 0.085 mmol) and 203 (499 mg, 0.22 mmol) were refluxed in toluene (10 ml) for 24 h, then the reaction solvent was removed under vacuum. Column chromatography on alumina was used to isolate the pure product, which was the first green band to be brought off using toluene as an eluent. The solvent was removed under vacuum and the product recrystallised by evaporation of a solution in 3:2 acetone/n-pentane at room temperature. Yield: 263 mg (62%). +ve electrospray: \textit{m/z}: 4939 [MH]+. \nu_{\text{max}} (\text{cm}^{-1}): 2029 (\text{C=O}), 1931 (\text{C=O}), 1890 (\text{C=O}). Elemental analysis calcd. (%) for C\textsubscript{177}H\textsubscript{310}ClCr\textsubscript{14}F\textsubscript{16}N\textsubscript{4}Ni\textsubscript{2}O\textsubscript{67}Re: C 43.06, H 6.33, N 1.13, Cr 14.74, Re 3.77. Found: C 42.73 H 6.32, N 1.07, Cr 14.36, Re 3.85.
ReCl(CO)$_5$([Pr$_2$NH$_2$][Cr$_7$CoF$_8$(O$_2$CCMe$_3$)$_{15}$(O$_2$CPy)])$_2$ 221

This compound was prepared and purified in manner similar to 222 by using ReCl(CO)$_5$ (20.0 mg, 0.055 mmol) and 207 (303 mg, 0.131 mmol), giving a green solid. Recrystallisation was achieved as for 222 using a 3:2 acetone/n-pentane mixture at room temperature. Yield: 87.7 mg (35%).

$\nu_{\text{max}}$ (cm$^{-1}$): 2021 (C=O), 1930 (C=O), 1887 (C=O).

Elemental analysis calcd. (%) for C$_{177}$H$_{310}$ClCo$_2$Cr$_{14}$F$_{16}$N$_4$O$_6$: C 43.05, H 6.33, N 1.23, Cr 14.74, Re 3.77. Found: C 43.00, H 6.62, N 1.04, Cr 14.54, Re 3.79.

[Re{[Pr$_2$NH$_2$][Cr$_7$NiF$_8$(O$_2$CCMe$_3$)$_{15}$(O$_2$CPy)]}(CO)$_3$](bpy)[PF$_6$] 222

To a solution of $\text{fac-ReCl(CO)}_3$(bpy) (0.106 g, 0.230 mmol) and 203 (2.53 g, 1.09 mmol) in refluxing toluene (55 ml) was added a solution of AgBF$_4$ (54 mg, 0.276 mmol) in toluene (5 ml). The reaction mixture was refluxed for 3 h then filtered warm to remove AgCl before the filtrate was refluxed for a further 23 h. The warm solution was filtered once more and the solvent was removed from the filtrate under vacuum. The product was extracted into acetonitrile (200 ml), which was then removed under vacuum. The solid was dissolved in a minimal amount of THF and metathesis to the PF$_6^-$ salt was effected by addition of aqueous NH$_4$PF$_6$. The green product was collected by filtration, washed with a large amount of water and dried under vacuum. Yield: 186 mg (28%). $^+$ve electrospray: $m/z$: 2915 [MNa]$^+$, 2742 [M – PF$_6$]$^-$, 2337 [{Pr$_2$NH$_2$} {Cr$_7$NiF$_8$(O$_2$CCMe$_3$)$_{15}$(O$_2$CPy)}Na]$^+$, 2315 [{Pr$_2$NH$_2$} {Cr$_7$NiF$_8$(O$_2$CCMe$_3$)$_{15}$(O$_2$CPy)}]$^+$. $\nu_{\text{max}}$ (cm$^{-1}$): 2037 (C=O) (s), 1926 (C=O) (br). Elemental analysis calcd. (%) for C$_{100}$H$_{165}$Cr$_{14}$F$_{14}$NaNiO$_{67}$PRe: C 41.57, H 5.76, N 1.94, Cr 12.60. Found: C 41.48, H 5.78, N 1.73, Cr 12.39.

[Ru(bpy)$_2$][Pr$_2$NH$_2$][Cr$_7$NiF$_8$(O$_2$CCMe$_3$)$_{15}$(O$_2$CPy))]$_2$][PF$_6$]$^-$ 223

cis-Ru(bpy)$_2$Cl$_2$•2H$_2$O (52 mg, 0.108 mmol) and AgCF$_3$SO$_3$ (87 mg, 0.339 mmol) were refluxed in ethanol (25 ml) for 3 h, giving a deep red solution. The solution was filtered warm to remove AgCl before 203 (1.174 g, 0.507 mmol) was added and further refluxing for 16.5 h produced an orange-green solution. The solution was cooled and filtered and the solvent was removed under vacuum. The product was extracted into acetonitrile, removing the starting wheel, and to the solution was added aqueous NH$_4$PF$_6$ to form an orange-brown precipitate, which was collected by filtration, washed with water and dried under
vacuum. Yield: 174 mg (32%). +ve electrospray: m/z: 2522 [M – 2PF₆]²⁺, 5193 [M – PF₆]⁺. λ_max (ε/mol⁻¹ dm³ cm⁻¹): 433 (12300), 367 (15600), 291 (67200). Elemental analysis calcd. (%) for C₁₉₄H₃₂₆Cr₁₄Ni₇F₂₈N₈O₆₄P₂Ru•1.5NH₄PF₆: C 41.76, H 6.00, N 2.38, Cr 13.05. Found: C 41.57, H 5.90, N 2.34, Cr 12.98.

\[ \text{[Cu(NO}_3\text{)}_2(H}_2\text{O)}(\text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_14(\text{O}_2\text{CPy})_2)]_n \] 224

To a warm solution of 208 (107 mg, 0.046 mmol) in acetone was added Cu(NO₃)₂•2.5H₂O (13.2 mg, 0.057 mmol) in acetone (10 ml) and the solution was refluxed for 10 min. During this time a green precipitate began to form. The reaction was allowed to cool to room temperature and left covered for 24 h, affording a precipitate. This solid was collected by filtration, washed with acetone and dried under vacuum. X-ray quality crystals were grown by evaporation of a tetrahydrofuran/toluene solution. Yield: 114 mg.
2.3 RESULTS AND DISCUSSION

2.3.1 Synthetic Studies

2.3.1.1 Substitution into Wheels

The primary starting materials, $[\text{Pr}_2\text{NH}_2][\text{Cr}_7\text{MF}_8(\text{O}_2\text{CCMe}_3)_{16}]$ where $M = \text{Ni (201)}$ and $\text{Co (202)}$ were synthesised following published procedures.$^5$ The cobalt analogue, however, required additional purification by washing the product through a silica column with toluene to remove impurities prior to recrystallisation.

Functionalisation of the wheels was achieved following a procedure used with the previously published ISNA wheel $^{203}$ (Scheme 1). Refluxing a roughly four-fold excess of a second carboxylic acid with 201 or 202 in propan-1-ol allowed this acid to be incorporated into the wheel. In order to aid the removal of the anionic pivalate ligands a polar solvent is required, but the solubility of the wheels must also be considered. Consequently, propan-1-ol was considered to be the most appropriate solvent for this reaction.

$$\begin{align*}
\text{LH, reflux propan-1-ol} & \quad \text{20 h} \\
[\text{Pr}_2\text{NH}_2][\text{Cr}_7\text{MF}_8(\text{O}_2\text{CCMe}_3)_{16}] & \quad [\text{Pr}_2\text{NH}_2][\text{Cr}_7\text{MF}_8(\text{O}_2\text{CCMe}_3)_{15}(L)]
\end{align*}$$

Scheme 1. Synthesis of substituted wheels.

$M = \text{Ni, } L =$

- 203
- 204
- 205
- 206
- 207

$M = \text{Co, } L =$
Isolation of the product was achieved by column chromatography on silica gel, eluting initially with toluene to remove the starting material, before increasing the gradient of the eluent using a 40–60 °C petroleum ether/ethyl acetate mixture (2 L:100 ml). Once the band of the mono-substituted product, \([\text{Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CMe}_3)_{15}(\text{L})]\), had separated from bands of more highly substituted products, the amount of ethyl acetate was increased to elute the desired product faster.

For certain substitution reactions, it is also possible to obtain disubstituted products (Scheme 2), although the yields of these are significantly lower than those of the comparable mono-substituted wheels. These minor products were brought off the same silica columns used to isolate the mono-substituted wheels simply by increasing the relative ethyl acetate content of the eluent.

\[
\begin{align*}
[\text{Pr}_2\text{NH}_2][\text{Cr}_7\text{MF}_8(\text{O}_2\text{CMe}_3)_{16}] & \xrightarrow{\text{LH, reflux propan-1-ol}} [\text{Pr}_2\text{NH}_2][\text{Cr}_7\text{MF}_8(\text{O}_2\text{CMe}_3)_{14}(\text{L})_2] \\
20 & \text{h}
\end{align*}
\]

Scheme 2. Synthesis of disubstituted wheels.

Substitution of the carboxylate occurs across the divalent metal and one of the \(\text{Cr}^{\text{III}}\) ions, because the former is much more kinetically labile. The substitution takes place in the axial position due to the trans effect of the opposite \(\pi\)-accepting carboxylate ligands. The equatorial pivalate groups are trans to \(\pi\)-donating fluoride ligands which strengthens their bonds to the metal centre. The second substitution also occurs on the divalent metal ion and an adjacent \(\text{Cr}^{\text{III}}\) ion. As it is preferential for the substitution to occur in the axial position, it occurs on the \(\text{Cr}^{\text{III}}\) ion that has not already been involved in the first substitution.
Table 2. Yields of substitution reactions.

<table>
<thead>
<tr>
<th>Product</th>
<th>Yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>203</td>
<td>32</td>
</tr>
<tr>
<td>204</td>
<td>12</td>
</tr>
<tr>
<td>205</td>
<td>6</td>
</tr>
<tr>
<td>206</td>
<td>44</td>
</tr>
<tr>
<td>207</td>
<td>24</td>
</tr>
<tr>
<td>208</td>
<td>3</td>
</tr>
<tr>
<td>209</td>
<td>6</td>
</tr>
<tr>
<td>210</td>
<td>3</td>
</tr>
</tbody>
</table>

The efficiency of the substitution reactions appears to vary widely (see Table 2), which has been attributed to a number of factors. Firstly, and most obviously, the solubility of the substituting carboxylic acid in propan-1-ol plays a major role as none of the carboxylic acids used are fully soluble, even at reflux. Another potentially important factor is the $pK_a$ of the incoming acid. The lower this $pK_a$, the higher the anticipated yield, due to the increased ability of the acid to dissociate into the anionic species required to substitute into the wheel.

2.3.1.2 Reactions of Substituted Wheels with First Row Transition Metals

Initial coordination chemistry of these novel functionalised wheels was attempted with first row transition metal centres. Due to the relative weakness of the bonds formed, coordination to first row d-block metals can generally occur readily, without the requirements of harsh conditions or much purification. It is not anticipated that selective control of magnetic interactions will be possible with these linked-wheel systems. However, preliminary work with the previously synthesised substituted wheel, 203, had already produced some interesting results, so this approach provides a good basis to compare the effectiveness of the various substituents in terms of metal ion coordination.

Systems of the type $\text{Cu}_2(\text{O}_2\text{CCMe}_3)_{4}[[\text{Pr}_2\text{NH}_2]_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{L})_2]_2$ were formed by reaction of the mono-substituted wheels with $\text{Cu}_2(\text{O}_2\text{CCMe}_3)_{4}(\text{HO}_2\text{CCMe}_3)_2$ in toluene. Stirring the reaction at room temperature and then at reflux produced the desired linked-wheel compounds and slow evaporation of the solvent afforded crystals. The wheels 204, 205 and 206 formed 211, 212 and 213, respectively (Scheme 3).
Crystals suitable for X-ray diffraction were obtained for the products 211 and 213, but those of 212 did not diffract strongly enough. Hence, 212 has been characterised by elemental analysis only. 211 and 213 have been characterised by elemental analysis and X-ray crystallography.

Following the experimental procedures used in the synthesis of the ISNA analogue 214, 204 was linked through a single copper centre to form 215 (Scheme 4). An acetone solution of Cu(NO₃)₂·2.5H₂O and 204 was heated for 5 min and then left to stand covered for 2 days forming crystals, which have been characterised by X-ray crystallography and elemental analyses. When a similar reaction was attempted with 206 however, crystals were not obtained. Therefore, the solution was allowed to stand only partially covered, producing some crystalline material. However, elemental analyses showed only trace quantities of copper, indicating that this material was unreacted 206.

Scheme 3. Synthesis of Cu dimer linked-wheel systems.

Scheme 4. Synthesis of copper nitrate linked wheel systems.
Having encountered difficulties in attempting to synthesise a linked-wheel system previously using 206 as a ligand, it was decided to investigate its general suitability as a ligand, using some alternative metal centres of type trans-M(hfac)$_2$(H$_2$O)$_2$ where M = Cu, Ni or Mn (Scheme 5). In a similar fashion to other reactions with first row neutral metal complexes, a 2:1 ratio of wheel to metal complex was mixed and allowed to crystallise, using the insolubility of the linked-wheel system to drive its formation. The first complex to be considered was the copper one, as it was with another copper complex where the problems had been discovered initially. In a similar fashion to the reaction described previously, no crystalline material was produced, so the solution was allowed to partially evaporate. However, elemental analyses on the material produced showed no copper to be present, so no link had been made. In stark contrast, use of 203 as a ligand afforded a crystalline material, 216, within five minutes.

![Diagram of linked-wheel system](attachment:linked_wheel_system.png)

**Scheme 5.** Synthesis of trans-M(hfac)$_2$(L)$_2$ (M = Cu, Ni or Mn) complexes.

Reacting 206 with trans-Ni(hfac)$_2$(H$_2$O)$_2$ in warm acetone for 5 mins afforded 217 in a 62% isolated yield. 217 was characterised by X-ray crystallography and elemental analyses, showing that 206 can be used as a monodentate ligand with nickel centres. Reaction of 206 with trans-Mn(hfac)$_2$(H$_2$O)$_2$ also appeared to be successful. However, the complexation occurred much more slowly when compared with the analogous Ni system, taking 5 days. As yet, crystals of 218 suitable for x-ray diffraction have not been produced.

It is unknown why 206 does not appear capable of coordinating in a monodentate fashion to a single copper centre, when it does form linked-wheel systems through other metal...
centres, especially as there is previously published evidence of 4-pyridazine being used as a monodentate ligand with copper. However, 4-pyridazine is an electron deficient ligand with the ortho-nitrogen removing electron density from the coordinating nitrogen, which may have an impact on its coordinating abilities.

A tri-wheel system was synthesised by reacting 3 equivalents of 204 with Fe₂NiO(O₂CCMe₃)₆ (HO₂CCMe₃)₃ in n-hexane at room temperature (Scheme 6). Crystallisation of the linked-wheel system formed (219) was attempted by many methods, but slow evaporation of a diethyl ether/acetone solution gave the best quality crystals, confirming the coordination of three wheels. However, satisfactory elemental analyses are still required before pursuing magnetic and EPR studies on this product. A similar reaction has also been attempted with 203, but there has been no evidence to date of successful synthesis of a linked-wheel system.

![Scheme 6. Synthesis of 219.](image)

2.3.1.3 Reactions with Second and Third Row Transition Metals

Reactions with first row transition metals occur very readily, allowing mild conditions and (often) facile crystallisation directly from the reaction solution. However, the coordination of second and third row transition metal centres is expected to provide more interesting systems from the perspective of redox/photoactivity, and also more stable products. The only substituted wheels considered in this section are the ISNA derivatives 203 and 207.

By reacting 203 with ReCl(CO)₅ in refluxing toluene for 24 h, it is possible to form the linked-wheel system 220 (Scheme 7). Purification was achieved on an alumina column.
using toluene as the eluent. An analogous reaction using 207 in place of 203 produced 221, but in a yield only half of that obtained for 222. Both of these products have been characterised using IR spectroscopy and elemental analyses, and the structure of 220 has been proven using X-ray crystallography.

Although it is possible to form linked-wheel systems from reactions with ReCl(CO)$_5$, such species are not expected to demonstrate any potentially useful photo/redox-activity. One method of introducing photoactivity into the system is to include a relatively low energy MLCT chromophore, using a co-ligand such as 2,2’-bipyridine. Consequently, it was decided to react 203 with the known complex fac-ReCl(CO)$_3$(bpy) to form 222. It was expected that only one wheel would coordinate to the Re centre due to selective substitution of the chloride ligand. However, it might be possible to use a complex such as fac,fac-{(ReCl(CO)$_3$)$_2$(bpym)} (bpym = 2,2’-bipyrimidine) to link two wheels.

222 was synthesised using a previously unpublished one-step method, recommended by Dr Mike Coogan of the University of Cardiff (Scheme 8). 1 equivalent of fac-
ReCl(CO)$_3$(bpy) was refluxed with 1.2 equivalents of AgBF$_4$ and 5 equivalents of 203 in toluene. After 3 h, the reaction mixture was filtered to remove AgCl, and the filtrate was refluxed for a further 22 h. Purification was achieved by metathesis to the PF$_6^-$ salt, followed by chromatography on an alumina column. Attempts to grow diffraction-quality crystals via slow evaporation of an acetone/methanol solution proved unsuccessful, but 222 has been characterised by elemental analyses, mass spectrometry and IR spectroscopy.

The reaction of 203 with cis-Ru(bpy)$_2$Cl$_2$•2H$_2$O produced 223 in ca. 30% yield, via a two-step method based on a literature procedure, that involves AgCF$_3$SO$_3$ in ethanol to generate the solvento intermediate cis-[Ru(bpy)$_2$(EtOH)$_2$][CF$_3$SO$_3$]$_2$ (Scheme 9). Again, no crystals have been grown, but characterisation of 223 has been achieved using mass spectrometry, UV/vis spectroscopy, cyclic voltammetry and elemental analyses.

2.3.1.4 Formation of polymers

A polymer of structure \{Cu(NO$_3$)$_2$(H$_2$O)([Pr$_2$NH$_2$][Cr$_7$NiF$_8$(O$_2$CCMe$_3$)$_{14}$(O$_2$CCpy)$_2$])$_2\}_n, 224, was formed by reacting the disubstituted wheel 208 with Cu(NO$_3$)$_2$•2.5H$_2$O (Scheme 10). Diffraction-quality crystals were obtained by slow evaporation of a THF/toluene solution.
2.3.2 Mass Spectrometry Studies

Mass spectrometry (MS) is a technique that has proved quite useful in the identification of “wheel” systems, as NMR spectroscopy cannot be used in general due to the fast relaxation of the Cr\text{III} nuclei contained in the backbones of the wheels. Previous investigations on single wheels had deduced that the best technique to observe the presence of the wheels is electrospray MS in THF, due to solubility factors. The ISNA-substituted wheel, 203, shows peaks for the [MH]\text{+} and [MNa]\text{+} ions in the positive electrospray spectrum. In the negative electrospray spectrum a peak is observed for the [M – Pr$_2$NH$_2$]\text{+} (wheel without template) ion. Each of the novel monosubstituted and disubstituted wheels, 204–210, has been characterised by MS, all showing the anticipated peaks.

The unusually high molecular weights are an issue when analysing linked-wheel systems. Each wheel has a molecular weight of approximately 2300 Da and the electrospray MS machine used at The University of Manchester can only measure $m/z$ to a maximum value of 3000, so the [M]\text{+} peaks cannot be observed for linked-wheel systems. However, analysis of 223 showed that MS is still a viable technique because a peak can seen quite clearly at $m/z = 2523$, which is associated with the [M – 2PF$_6$]\text{2+} dication. Simulation of this peak pattern (Figure 40) shows agreement with the observed spectrum.
Later on in the project a collaboration was commenced with Dr Lindsay Harding of the University of Huddersfield who offered to study some of the linked-wheel systems using an ESI-TOF machine, which has a wider mass range. Two linked green wheel systems were analysed successfully, 220 and 223. The spectrum of 220 shows a peak for [MH]$^+$ at $m/z = 4939$, and a peak is seen at $m/z = 5193$ in the spectrum of 223 corresponding to [M – PF$_6$]$^+$. The mass peak of the three-wheel system, 219, however is not observed as its molecular weight of 7818 Da appears to be too high for even this technique.

### 2.3.3 Infrared Spectroscopy

Infrared spectra have been recorded for the three Re-based compounds, 220, 221 and 222. The spectra of 220 and 221 are virtually identical, which is as expected as the only difference between these two compounds is the divalent metal contained in the wheels. Three ν(CO) bands can be seen between 2030 and 1885 cm$^{-1}$, as expected for fac-Re(CO)$_3$ complex fragments. As can be seen in Table 3, these bands occur at energies similar to those reported for the model complex fac-ReCl(CO)$_3$(py)$_2$, indicating that the presence of the wheels does not have a significant electronic effect on the Re centre. In a similar fashion, the bands associated with the wheel ligands are not shifted upon coordination to the Re centre, as shown by direct comparison of the spectra for 203 and 220 (Figure 41).
Table 3. IR data for novel and reference Re complexes in the CO stretching region.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Bands, cm$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$fac$-$\text{ReCl(CO)}_3(\text{py})_2$</td>
<td>2025 1920 1885</td>
</tr>
<tr>
<td>220</td>
<td>2029 1931 1890</td>
</tr>
<tr>
<td>221</td>
<td>2021 1930 1887</td>
</tr>
<tr>
<td>222</td>
<td>2037 1926 (br)</td>
</tr>
<tr>
<td>$fac$-$[\text{Re(py)(CO)}_3(bpy)]^{1104}$</td>
<td>2026 1906 (br)</td>
</tr>
</tbody>
</table>

Figure 41. IR Spectra of 203 and 220.

Figure 42. IR Spectrum of 222.
The IR spectrum of 222 (Figure 42) shows only two $\nu$(CO) bands. The two lower energy bands, instead of being well defined, are merged into one broad band which is a common observation for complexes containing $[\text{ReL(CO)}_3(\text{bpy})]^{+}$ fragments.\(^{95}\)

### 2.3.5 UV/Visible Spectroscopy

![UV/vis spectrum of 223 in acetonitrile.](image)

The UV/vis spectrum of 223 can be seen in Figure 43. As expected, two $d\rightarrow\pi^*$ MLCT bands are observed, which are not dissimilar from the literature data obtained for cis-$[\text{Ru(bpy)}_2(\text{py})_2][\text{PF}_6]_2$, helping to confirm that the expected product has been formed.\(^{96}\) A shoulder is observed at approximately 500 nm, which could be attributable to $d\rightarrow\pi^*$ interactions between the Ru centre and the wheel ligand.

### 2.3.6 Electrochemistry

A cyclic voltammogram of 223 was run in a 0.1 M $[\text{NBu}^n_4][\text{PF}_6]$ solution in acetonitrile, and a representative trace is shown in Figure 44. The trace is very weak due to the high molecular mass of the product as well as its poor solubility in acetonitrile. What appears to be a reversible Ru$^{III/II}$ oxidation can be seen at 1.43 V vs. Ag-AgCl ($\Delta E_p = 55$ mV), a potential 0.13 V higher than that observed for cis-$[\text{Ru(bpy)}_2(\text{py})_2][\text{PF}_6]_2$,\(^{97}\) implying a
greater stability of the Ru\textsuperscript{II} oxidation state. Three irreversible waves can also be seen at -1.27 V, -1.35 V and -1.60 V vs. Ag-AgCl related to ligand reductions.

![Cyclic voltammogram of 223 in acetonitrile at a scan rate of 200 mV s\textsuperscript{-1}.](image)

**Figure 44.** Cyclic voltammogram of 223 in acetonitrile at a scan rate of 200 mV s\textsuperscript{-1}.

### 2.3.7 EPR Studies

The EPR spectra of the copper dimer linked-wheel systems 211 and 213 have been recorded. In a fashion similar to the previously published ISNA system\textsuperscript{65} these spectra show no changes when compared with that of the unlinked wheel (Figure 45), with a simple $S = \frac{1}{2}$ signal. It appears that the strongly antiferromagnetically coupled copper dimer moiety prevents coupling between the wheels. The Re linked-wheel system 221 shows an essentially identical spectrum.

![Q-band EPR spectrum of an individual wheel at 5 K.](image)

**Figure 45.** Q-band EPR spectrum of an individual wheel at 5 K.
However, the EPR spectrum of 215 differs significantly from that of the individual wheel. The spectrum of 215 together with that of the previously published compound 214 can be seen in Figure 46. While the parameters have not yet been established for 215 it is anticipated that the interaction between the wheels will be weaker than for 214.

![Figure 46. Q-band EPR spectra of 214 (red) and 215 (black).](image)

The EPR spectrum of the copper linked polymeric chain of wheels was also run, giving a broad spectrum. The spectrum is too broad to be modelled, but differs from the unlinked wheels hence interaction of some sort is occurring.

![Figure 47. Q-band EPR spectrum of 224.](image)
2.3.9 X-Ray Crystallography

Due to the magnetic nature and large size of these linked-wheel systems, it is very difficult to confirm their structures without using X-ray crystallography. Many of these systems have therefore been characterised by this method.

Crystal structures of 206, 207 and 210 have been determined. However, these structures have not been fully solved due to pronounced disorder of the divalent metal and hence the substituted carboxylate. Also, the mass spectra for these compounds confirm the formation of the desired structures, showing an instantaneously recognisable pattern, so having only low-quality X-ray structures is sufficient for our purposes.

For all of the compounds discussed in this chapter, the structures of the wheels are consistent, and agree with previously published structures. The Cr–F bond lengths average between 1.90 and 1.92 Å and the Cr–O bond lengths are between 1.94 and 1.99 Å. The Ni–F bond lengths are slightly longer, between 2.00 and 2.02 Å with the Ni–O bond lengths averaging at 1.99–2.01 Å. In all cases the [Pr₂NH₂]⁺ template can be seen in the central cavity of all the wheels, held in place by N–H···F bonds. The only differences observed are the carboxylates that are substituted into the wheels.

Figure 48. A representation of the molecular structure of 211.

Structures have been obtained for 211 (Figure 48) and 213 (Figure 49), from crystals formed directly from their reaction mixtures in toluene. Both form monoclinic systems, with 211 adopting a space group C₁2/C₁ and 213 P₁2₁/C₁. Both structures show identical
central regions with a Cu paddlewheel dimer linking the wheels together. The only significant difference between the two central portions is the Cu–N bond distances, with that in 211 being ca. 0.03–0.04 Å shorter than that in 213. This observation might explain the poor yield obtained for 213, with a weaker bond being formed between the pyridazine ring and Cu centre. The disordered nature of the six-membered rings precludes any meaningful comparisons of their geometric parameters. The Cu–N bond in 213 (2.183(5) Å) is also significantly longer than that in a monodentate pyridazine Cu complex found in the literature90 (1.984(2) Å), suggesting that the attached wheel may weaken the effectiveness of 209 as a ligand.

Figure 49. A representation of the molecular structure of 213.

Measurable crystals of 215 were grown upon formation of the product in an acetone solution. The crystals were twinned, and thus a poor final $R_1$-factor of 0.1789 was obtained. A picture of the structure can be seen in Figure 50. The crystals are triclinic in the $P\bar{1}$ space group. The wheels are linked in a virtually trans arrangement, with an N–Cu–N angle of 172.0(3)º, through a five-coordinate Cu centre. Also coordinated are two monodentate nitrate ligands and an acetone solvent molecule, which demonstrates Jahn-Teller distortion with a Cu–O bond distance of 2.431(6) Å as opposed to the shorter nitrate Cu–O bonds which have an average length of 1.975 Å. The two nitrate groups are apparently coordinated in a monodentate fashion, due to a significant difference in the Cu–O distances (0.65 Å and 0.77 Å). However, 0.77 Å is still shorter than the combined values of the Van der Waals radii of the Cu and O atoms, so it is plausible that at least one
of the nitrate ligands could be bidentate. The arrangements of the other Cu-ligand bonds, which are all close to 90°, support the idea of an octahedral structure.

Figure 50. A representation of the molecular structure of 215.

Figure 51. A representation of the molecular structure of 217.

In a similar fashion to the crystal structures mentioned thus far, crystals of 217 were obtained directly from the reaction to produce this compound in acetone. This structure is orthorhombic in the Pccn space group, and has the wheels coordinated in a cis geometry (Figure 51), with an N–Cu–N angle of 89.9(4)°. In contrast, the ISNA analogue shows a
trans disposition of the wheels.\textsuperscript{98} As in the structure of 213, the ortho nitrogens in 217 are disordered over two sites.

![Figure 52](image_url) A representation of the molecular structure of 220.

Crystals of 220 were grown by evaporation of an $n$-pentane/acetone solution. The crystals are monoclinic in a $P2_1/c$ space group and the molecular structure can be seen in Figure 52. Unfortunately, due to disorder about the Re centre the $R_1$ factor is very high at 14%. However, it is clear that the desired compound has been formed. The Re atom is disordered over two sites, with N–Re–N angles of 86–88°.

The crystal structure of 219 adopts the triclinic $P\overline{1}$ space group. This complex (Figure 53) contains three wheels coordinated through 4pyac to the central triangular metal cluster. The central oxygen is positioned almost symmetrically, with the Ni–O bond being only slightly longer (~0.02 Å) than the Fe–O bonds. The three metal centres in the triangle have very slightly distorted octahedral geometries, with bond angles in the range 82–97°.
The final structure to be considered is that of 224, crystals of which were grown by evaporation of a THF/toluene solution at room temperature. 224 adopts the monoclinic space group $P2_1/n$, and the polymeric complex (Figure 54) contains coordinated THF derived from the substitution of water during crystallisation. As with the monomeric
structure, 215, the Cu–O bond to this solvent molecule (2.249 (12) Å) is between 0.23 and 0.41 Å longer than the other Cu–O bonds. The structure of the disubstituted wheel is clearly resolved and as observed previously,98 with both ISNA groups coordinated to the divalent metal centre and an adjacent CrIII ion.
2.4 CONCLUSIONS

The series of mono- and disubstituted wheels has been extended with varying degrees of success. These novel wheels have been reacted with a variety of 1st row transition metal complexes, in order to assess their suitability as ligands. 204 appears to behave well in complexation reactions, but 206 has not been coordinated to single Cu metal centres. However, crystal structures have been obtained for the 206-containing complexes 213 and 217, proving that it can act as a ligand in certain situations. EPR studies have been carried out on the 204-containing system 215, and comparison with the 203-containing 214 demonstrates that the interaction between the wheels weakens as the link extends on moving from ISNA to 4-pyridylacrylate.

Some further work has been undertaken with 203 as a ligand for 2nd and 3rd row Re and Ru complexes expected to form MLCT/redox-active complexes. An X-ray crystal structure has been obtained for 220, confirming the coordination to the Re centre. While no crystal structure has of yet been obtained for 223, the data available indicate that the desired product has been formed and that it shows MLCT absorption bands and a fully reversible Ru$^{2+/3+}$ oxidation, suggesting that switchability may be possible with this system.

Reaction of a bis-ISNA-substituted wheel with Cu(NO$_3$)$_2$•2.5H$_2$O produces a polymeric structure, which has been fully characterised. The EPR spectrum shows some evidence of interaction occurring between the wheels, but the spectrum is too broad to be able to model any parameters.

2.6 FURTHER WORK

While a crystal structure has been solved for 220, a better structure will be required for publication purposes. The use of different solvent systems could be attempted in order to grow crystals of higher quality, or the existing crystals could be sent to a synchrotron in order to get better data. Obtaining a crystal structure of 223, which is a good candidate for switching studies, is an important objective in order to fully characterise this product. This complex is extremely soluble as its PF$_6^-$ salt, and consequently anion metathesis may be the best option to allow effective crystallisation. Metathesis to a bulkier anion such as
BPh$_4^-$ may improve the potential for packing in a lattice, as the complex itself is relatively large. Once a hypothetically switchable system containing two wheels has been fully characterised, then physical studies to assess the possibilities for potential qubit behaviour will be pursued.

Further substitution reactions into the wheels could be attempted using polypyridyl carboxylic acids, such as those shown in Figure 55. These could produce excellent ligands for coordination to metal centres such as Ru$^{II}$ or Re$^I$, leading to the possibility of having low energy MLCT excitations directed at the wheels themselves.

![Figure 55. Examples of polypyridyl carboxylic acids, potentially suitable for substitution into the wheels.](image)

While initial studies have involved attaching 203 to Re cores, using as precursors ReCl(CO)$_3$ and fac-[ReCl(CO)$_3$(bipy)]PF$_6$, other Re complexes may be considered in order to access systems that show low energy MLCT bands and also contain multiple wheels. One such complex precursor is {fac-ReCl(CO)$_3$}$_2$(bpym), which should be amenable to chloride substitutions. If this approach were to prove successful, then the same chemistry should be extendable to include larger polynuclear Re systems, such as triangles and squares. 99
Chapter 3 – Linked Purple Wheel Systems
3.1 INTRODUCTION

Purple wheels were first synthesised when experimenting with altering the templating amine molecule to observe what effect it would have on the size and shape of the resultant complex. Purple wheels differ from their green counterparts, in that the “template” used to form the wheel, in this case a sugared amine, \(N\)-ethyl-D-glucamine, coordinates directly into the wheel, replacing five of the eight fluorides. The purple derivatives also contain one less carboxylate ligand, allowing the template to coordinate twice to the chromium adjacent to the divalent metal, leaving a spare site at the divalent metal. In the uncoordinated purple wheel a solvent molecule can be found at this site, but this can be replaced easily by a good coordinating group such as a pyridine.

![Figure 56. Purple wheels linked through 4,4'-bpy.](image)

A reaction was attempted using 4,4'-bpy as the coordinating ligand, forming a linked-wheel system (Figure 56).\(^60\) Other reactions using the slightly longer linker bpe and molecules with more than two pendant pyridyl groups also proved successful. Applying this principle opens the door to systems linked by not only other organic molecules but also transition metal complexes that contain ligands with terminal pyridyl groups such as quaterpyridine (qpy) and its derivatives. By using such complexes it may be possible to
construct switchable systems. Rhenium(I) and ruthenium(II) complexes often display long-lived MLCT excited states and ruthenium(II) and iron(II) complexes undergo reversible oxidations.

Various organics and metal complexes (Figure 57) have been identified as suitable potential target linker molecules; some of these are already known in the literature, while others are novel. The work covered in this chapter includes the further development of novel linked-wheel systems with organic bridges, as well as studies towards the creation of a series of transition metal polypyridyl linked-wheel complexes.

![Figure 57. Target compounds for linking purple wheels together.](image-url)
3.2 EXPERIMENTAL PROCEDURES

3.2.1 Materials and Procedures

The compounds \((E,E,E)-1,6\text{-bis}(4\text{-pyridyl})-1,3,5\text{-hexatriene} \text{ (bph)}\),\(^{100}\) \(1,4\text{-bis}[E-2-(4\text{-pyridyl})\text{ethenyl}]\text{benzene} \text{ (bpv)}\),\(^{101}\) \(2,2':4,4':4',4'''\text{-quaterpyridyl} \text{ (qpy)}\),\(^{102,103}\) \(4,4'\text{-bis-[(E)-2-(4-pyridyl)vinyl]-2,2'-bipyridyl} \text{ (bbpe)}\),\(^{104}\) \(4'-(4''\text{-pyridyl})-2,2':6'2''\text{-terpyridine} \text{ (pytpy)}\),\(^{105}\) \([\text{Fe(pytpy)}_2][\text{BF}_4]_2\),\(^{105}\) \([\text{Ru(pytpy)}_2][\text{PF}_6]_2\),\(^{106}\) \(\text{cis-}[\text{Ru(bipy)}_2(\text{qpy})][\text{PF}_6]_2\),\(^{107}\) \(\text{fac-}[\text{ReCl(CO)}_3(\text{qpy})]_6\)\(^{108}\) and \([\text{Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})(\text{H}_2\text{O})]^{60}\) were synthesised by literature methods. All other starting materials were purchased commercially and used as supplied.

3.2.2 Physical Measurements

All \(^1\text{H} \text{NMR} \) spectra were recorded on Bruker Ultrasound 500 MHz or 300 MHz spectrometers. IR data were recorded using a Varian Bio-Rad Excalibur series IR instrument and UV data were recorded on a Shimadzu UV-2401 PC UV-VIS spectrophotometer. Elemental analyses were performed by the Microanalytical Laboratory, University of Manchester. Mass spectra were obtained using electrospray analysis on a Micromass Platform spectrometer at The University of Manchester. Linked-wheel systems were sent to Dr Lindsay Harding at the University of Huddersfield for analysis, and spectra were run using a Bruker MicroTOF-Q mass spectrometer with an electrospray ionisation source. EPR studies were performed by Dr Floriana Tuna and John Machin at The University of Manchester. Q-Band EPR was run on an Elexsys E500 spectrometer equipped with a Bruker 2 tesla magnet. Variable temperature studies were achieved using an Oxford VT liquid helium cryostat between 5 K and 30 K.
3.2.3 Syntheses

\[\text{[(bph)}\{\text{[Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}\text{(EtGlu)}]}\}_2\] 301

A solution of bph (27.3 mg, 0.117 mmol) in dichloromethane (10 ml) was added to a solution of [Cr\(_7\)NiF\(_3\)(O\(_2\)CCMe\(_3\))\(_{15}\)(EtGlu)(H\(_2\)O)] (513 mg, 0.231 mmol) in dichloromethane (10 ml) and the reaction stirred at room temperature for 15 h. The solvent was removed under vacuum and the dry product was washed with acetone to remove any excess starting material, producing a purple solid. The product was then recrystallised by evaporation at room temperature of a dichloromethane/acetone solution. Yield: 318 mg (58%). Elemental analysis calcd. (%) for C\(_{182}\)H\(_{312}\)Cr\(_{14}\)F\(_6\)N\(_4\)Ni\(_2\)O\(_{70}\)•CH\(_2\)Cl\(_2\): C 46.56, H 6.70, N 1.19, Cr 15.42, Ni 2.49. Found: C 46.70, H 6.84, N 1.06, Cr 15.11, Ni 2.09.

\[\text{[(bpvb)}\{\text{[Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}\text{(EtGlu)}]}\}_2\] 302

This compound was prepared and purified in a manner similar to 301 by using bpvb in place of bph (34.4 mg, 0.121 mmol) and [Cr\(_7\)NiF\(_3\)(O\(_2\)CCMe\(_3\))\(_{15}\)(EtGlu)(H\(_2\)O)] (509 mg, 0.230 mmol) in dichloromethane (20 ml), forming a purple solid. Diffraction-quality crystals were obtained by evaporation of a tetrahydrofuran/acetonitrile solution. Yield: 208 mg (38%). +ve electrospray: 4687 [MH]+, 2485 [{M −[Cr\(_7\)NiF\(_3\)(O\(_2\)CCMe\(_3\))\(_{15}\)(EtGlu)}]}H]+. Elemental analysis calcd. (%) for C\(_{186}\)H\(_{314}\)Cr\(_{14}\)F\(_6\)N\(_4\)Ni\(_2\)O\(_{70}\)•CH\(_2\)Cl\(_2\): C 47.08, H 6.68, N 1.17, Cr 15.26, Ni 2.46. Found: C 47.15, H 6.71, N 1.15, Cr 15.26, Ni 2.56.

\[\text{[[Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}\text{(EtGlu)}]}\{(\text{qpy})\}\{\text{[Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}\text{(EtGlu)}]}\] 303

To a solution of [Cr\(_7\)NiF\(_3\)(O\(_2\)CCMe\(_3\))\(_{15}\)(EtGlu)(H\(_2\)O)] (610 mg, 0.274 mmol) in dichloromethane (15 ml) was added a solution of qpy (42.0 mg, 0.134 mmol) in dichloromethane (5 ml) and the reaction was stirred at room temperature for 30 mins. The solution was then heated to reflux for a further 10 mins. The solution was concentrated under vacuum until only approximately 5 ml remained, then acetone (25 ml)
was added and the resultant solution was refluxed for 1 h. A purple precipitate was formed during this time and upon cooling of the solution it was collected by filtration, washed with acetone and dried in the air. Yield: 336 mg (53%). Elemental analysis calcd. (%) for C_{269}H_{459}Cr_{21}F_{9}N_{7}Ni_{3}O_{105}: C 46.75, H 6.69, N 1.42, Cr 15.80, Ni 2.55. Found: C 47.09, H 6.55, N 1.67, Cr 15.46, Ni 2.55.

\[\text{[[[Cr}_7\text{NiF}_4(\text{O}_2\text{CCMe}_3)]_{14}(\text{EtGlu})]}(\text{bbpe})\text{[[Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)]_{15}(\text{EtGlu})]}\]

To a solution of \([\text{Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)]_{15}(\text{EtGlu})(\text{H}_2\text{O})]\) (503 mg, 0.226 mmol) in dichloromethane (10 ml) was added bbpe (29.0 mg, 0.080 mmol) in dichloromethane, and the reaction stirred at room temperature for 15 h. The solvent was removed under vacuum and an acetone/acetonitrile recrystallisation gave a purple solid. Yield: 58 mg (10%). Elemental analysis calcd. (%) for C_{273}H_{463}Cr_{21}F_{9}N_{7}Ni_{3}O_{105}: C 47.09, H 6.70, N 1.41, Cr 15.68, Ni 2.53. Found: C 46.94, H 6.80, N 1.40, Cr 15.37, Ni 2.51.

**fac-ReCl(CO)$_3$(bbpe) (attempted) 306**

A solution of \(\text{ReCl(CO)}_5\) (100 mg, 0.276 mmol) and bbpe (105 mg, 0.290 mmol) in a 3:1 mixture of THF and toluene (20 ml) was refluxed for 8 h. The solution was allowed to cool, then filtered and the insoluble orange solid washed with toluene. Yield: 152 mg (69%). \(v(\text{cm}^{-1})\): 2015 s (C=O), 1841 br (C=O). Elemental analysis calcd. (%) for C$_{27}$H$_{18}$ClN$_4$O$_3$Re$_{1/3}$ReCl(CO)$_5$: C 43.66, H 2.30, N 7.10, Re 31.48. Found: C 43.93, H 2.43, N 6.93, Re 28.56.

**cis-[Ru(2,2′-bpy)$_2$(bbpe)][PF$_6$]$_2$ 308**

A solution of \(\text{cis-RuCl}_2(2,2′\text{-bpy})_2\cdot 2\text{H}_2\text{O}\) (101 mg, 0.194 mmol) and bbpe (70.5 mg, 0.195 mmol) in ethylene glycol (20 ml) was heated at reflux under argon for 1 h. The bright orange-red solution was cooled to room temperature and filtered. Aqueous NH$_4$PF$_6$ was added to the filtrate yielding a red precipitate, which was collected by filtration and washed with water. The crude product was purified on a silica gel column eluting with 0.3 M NH$_4$PF$_6$ in 1:1 acetone/acetonitrile, affording the product as the 3rd fraction off the column. Purification of this fraction was achieved by evaporating to
Chapter 3 – Linked Purple Wheel Systems

dryness then redissolving in a minimal amount of acetonitrile and adding aqueous NH₄PF₆. The precipitate was filtered off, washed with water and dried. A final reprecipitation from acetone/diethyl ether, collection by filtration, washing with diethyl ether and air drying gave a dark red solid. Yield: 94 mg (35%). ¹H NMR (CD₃CN) δH/ppm: 7.45 (4 H, t, J = 5.6 Hz, bpy), 7.60 (4 H, d, J = 7.3 Hz, bpy), 7.76 (2 H, d, J = 5.5 Hz, bpy) 7.78–7.84 (8 H, bpy, CH=CH), 7.99 (4 H, d, J = 6.6 Hz, bpy), 8.11 (4 H, t, J = 7.8 Hz, bpy), 8.55 (4 H, d, J = 8.3 Hz, bpy), 8.70 (4 H, d, J = 6.6 Hz, bpy), 8.84 (2 H, s, bpy). +ve electrospray: m/z = 920 [(M – PF₆)⁺], 388 [(M – 2PF₆)²⁺]. λ_{max}/nm (ε/M⁻¹ cm⁻¹): 472 (24,500), 376 (18,800), 289 (105,300), 257 (45,600). Elemental analysis calcd. (%) for C₄₄H₃₄F₁₂N₈P₂Ru•2HPF₆: C 38.92, H 2.67, N 8.25, P 9.13. Found: C 39.95, H 2.61, N 8.34, P 8.02.

cis-[Fe(2,2’-bpy)₂(bbpe)][PF₆]₂ 311

A portion of bbpe (69.0 mg, 0.190 mmol) was heated in acetone (50 ml) until fully dissolved. To this solution was added 2,2’-bpy (60.0 mg, 0.384 mmol) in acetone (5 ml) and the solution heated to reflux. A solution of [Fe(H₂O)₆][BF₄]₂ (66.0 mg, 0.196 mmol) in acetone (5 ml) was added dropwise to the solution causing an instantaneous colour change to blood red. The solution was stirred and heated for a further 5 min then filtered and the solvent was removed under vacuum. Metathesis of the impure product to the PF₆⁻ salt was achieved by addition of aqueous NH₄PF₆ to a concentrated acetonitrile solution. Purification was achieved on a silica gel column, eluting initially with 0.1 M NH₄PF₆ in 1:1 acetone/acetonitrile to remove [Fe(2,2’-bpy)₃][PF₆]₂, then 0.1 M NH₄PF₆ in 4:1 acetone/acetonitrile to isolate the desired product. Yield: 74 mg. ¹H NMR ((CD₃)₂CO) δH/ppm: 7.62 (4 H, t, J = 6.2 Hz, bpy), 7.76–7.82 (6 H, bpy), 7.90 (2 H, d, J = 5.3 Hz, bpy), 7.94 (4 H, s, HC=CH), 8.00 (4 H, d, J = 4.0 Hz, bpy), 8.31 (4 H, t, J = 6.9 Hz, bpy), 8.88 (8 H, d, J = 6.9 Hz, bpy), 9.13 (2 H, s, bpy). λ_{max}/nm: 536, 410, 336, 300 nm. Elemental analysis calcd. (%) for C₄₄H₃₄FeF₁₂N₈P₂: C 51.76, H 3.36, N 10.98, P 6.07. Found: C 35.70, H 2.36, N 7.34, P 5.97.
cis-[Fe(2,2'-bpy)₂(qpy)][PF₆]₂ (attempted) 310

This compound was prepared in a fashion identical to 311 but using 2,2'-bpy (61.2 mg, 0.392 mmol) in acetone (5 ml), qpy (59.8 mg, 0.193 mmol) in acetone (45 ml) and [Fe(H₂O)]₆[BF₄]₂ (65.0 mg, 0.193 mmol) in acetone (5 ml) forming a blood red solid. Currently this has not been isolated in a pure form.

fac-[ReCl(CO)₃(qpy)]{[Cr₇NiF₃(O₂CCMe₃)₁₅(EtGlu)]₃} 312

fac-ReCl(CO)₃(qpy) (48.2 mg, 0.133 mmol) and [Cr₇NiF₃(O₂CCMe₃)₁₅(EtGlu)(H₂O)] (547 mg, 0.246 mmol) were dissolved in dichloromethane (25 ml) and the reaction stirred at room temperature in the dark for 25 h. The solvent was then removed under vacuum. The residue was dissolved in 2:1 acetone/acetonitrile (30 ml), stirred briefly and allowed to sit partially covered overnight to give a precipitate. Recrystallisation was effected by gentle evaporation of a dichloromethane/acetonitrile solution. Yield: 206 mg (32%). +ve electrospray: 2201 [{Cr₇NiF₃Piv₁₅EtGlu}H]⁺, 2224 [{Cr₇NiF₃Piv₁₅EtGlu}Na]⁺, 5020 [MH]⁺; -ve electrospray: 616 [ReCl(CO)₃(qpy)]⁻, 2199 [{Cr₇NiF₃Piv₁₅EtGlu} – H]⁻. ν(cm⁻¹): 2026 (C=O), 1933 (C=O), 1910 (C=O). λmax/nm (ε/M⁻¹ cm⁻¹): 556 (1,600), 419 (6,600), 308 (25,500). Elemental analysis calcd. (%) for C₁₈₉H₃₁₂ClCr₁₄F₈N₆Ni₂O₇₃Re •2CH₂Cl₂: C 43.86, H 6.09, N 1.57, Cr 14.21, Re 3.68.

cis-[Ru(2,2'-bpy)₂(qpy)]{[Cr₇NiF₃(O₂CCMe₃)₁₅(EtGlu)]₂}[PF₆]₂ 313

A solution of cis-[Ru(2,2'-bpy)₂(qpy)][PF₆]₂ (45.6 mg, 0.045 mmol) and [Cr₇NiF₃(O₂CCMe₃)₁₅(EtGlu)] (203 mg, 0.091 mmol) in acetone (20 ml) was stirred at room temperature for 18 h. The solution was filtered and the solvent removed under vacuum. The product was extracted into acetonitrile, the solution filtered and evaporated to dryness under vacuum. Extraction into toluene in a similar manner produced a purple-red powder. Yield: 145 mg (60%). λmax/nm (ε/M⁻¹ cm⁻¹): 463 (16,700), 360 (13,100), 288 (71,200), 250 (72,000). Elemental analysis calcd. (%) for C₂₀₆H₃₄₀Cr₁₄F₁₈N₁₀
Ni$_2$O$_{70}$P$_2$Ru•C$_7$H$_8$: C 46.35, H 6.36, N 2.54, Cr 13.19, Ni 2.13, P 1.12. Found: C 46.29, H 6.40, N 2.30, Cr 12.90, Ni 1.94, P 1.32.

\[ \textit{cis} - [\text{Ru}(2,2'\text{-bpy})_2(\text{bbpe})][\text{Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})]\text{]}_2][\text{PF}_6]_2 \text{ 314} \]

This compound was prepared and purified in a manner similar to 313 by using \textit{cis} - [Ru(2,2'-bpy)$_2$(bbpe)][PF$_6$]$_2$ (25.0 mg, 0.018 mmol) and [Cr$_7$NiF$_3$(O$_2$CCMe$_3$)$_{15}$(EtGlu)] (115 mg, 0.052 mmol) in acetone (15 ml), forming a purple-red solid. Yield: 56 mg. $\lambda_{\text{max}}$/nm ($\varepsilon$/M$^{-1}$ cm$^{-1}$): 478 (12,900), 386 (10,100), 289 (55,300). Elemental analysis calcd. (%) for C$_{210}$H$_{344}$Cr$_{14}$F$_{18}$Ni$_2$O$_{70}$P$_2$Ru: C 46.03, H 6.33, N 2.56, Cr 13.29, Ni 2.14, P 1.13. Found: C 45.47, H 6.40, N 1.91, Cr 13.01, Ni 2.06, P 1.43.

\[ \text{[Fe(pytpy)$_2$}[\text{Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})]\text{]}_2][\text{BF}_4]_2 \text{ 315} \]

This compound was prepared and purified in a manner similar to 313 by using [Fe(pytpy)$_2$][BF$_4$]$_2$ (68.7 mg, 0.081 mmol) and [Cr$_7$NiF$_3$(O$_2$CCMe$_3$)$_{15}$(EtGlu)] (503 mg, 0.227 mmol) in acetone (25 ml), forming a bright purple solid. Yield: 77 mg (18%). Elemental analysis calcd. (%) for C$_{206}$H$_{338}$B$_2$Cr$_{14}$F$_{14}$FeN$_{10}$Ni$_2$O$_{70}$: C 47.01, H 6.47, N 2.66, Cr 13.83, Ni 2.23. Found: C 47.25, H 6.33, N 2.43, Cr 13.29, Ni 2.15.
3.3 RESULTS AND DISCUSSION

3.3.1 Synthesis of Organic Linker Molecules and Ligands

All organic molecules and ligands used in this chapter (Figure 58) were synthesised via published experimental procedures; Dr Simon Foxon provided all of these compounds, with the exception of pytpy.

![Figure 58. Organic linker molecules.](image)

A slightly adapted method was used to synthesise pytpy (Scheme 11), due to problems encountered when trying to isolate the product of the first step, 1,5-bis(2-pyridyl)-3-(4′-pyridyl)pentane-1,5-dione. The literature method\(^{105}\) indicated that upon completing the first step of the reaction, the intermediate product may be formed as an orange gum and the addition of methanol would allow the isolation of a white solid. It was discovered that this approach did not work as the initially formed white solid almost instantaneously reverted to orange gum and the addition of further methanol redissolved the product. Many different solvent combinations, primarily methanol and water mixtures, were attempted to allow isolation of the white solid, but without success. Consequently, it was decided to attempt the second step of the reaction using the orange gum, isolated by decanting off the reaction solvent. This strategy worked, albeit in a less effective manner...
than published, and the pytpy was isolated in an 8.6% yield as off-white crystals, which were characterised by $^1$H NMR, mass spectrometry and elemental analysis.

![Scheme 11. Synthesis of pytpy.](image)

3.3.2 Synthesis of Organic Linked-wheel systems

![Scheme 12. Synthesis of two-wheel systems with organic linkers.](image)

In a fashion similar to the method used for the previously synthesised 4,4'-bipy and bpe linked-wheel systems,$^{60}$ the syntheses of 301 and 302 were achieved simply by reacting the two starting materials together at room temperature (Scheme 12). Isolation of the products was achieved by exploiting their differences in solubility when compared to both starting materials. Various attempts were made at crystallising 301 and 302. The crystallisation approach considered, as for most wheel systems, was evaporation of a mixed solvent system at room temperature. A THF/acetonitrile solution proved successful in producing diffraction-quality crystals of 302, but no measurable crystals have been obtained for 301 to date.
Chapter 3 – Linked Purple Wheel Systems

The compound qpy is an ideal ligand to link wheels through metal centres due to its free terminal pyridyl groups available upon chelation to metal centres. Prior to reacting the wheels with qpy complexes, the wheels were reacted with qpy alone. Using a 2.5:1 ratio of wheel to qpy, followed by gentle evaporation of a diethyl ether/acetone solution, produced an interesting crystal structure.

Instead of the anticipated two-wheel system, a three-wheel system had formed (Scheme 13). This was not wholly unexpected as qpy has 4 free nitrogen atoms available for coordination to the metal wheels. What is unexpected is that the two adjacent nitrogens allow bidentate coordination of the third wheel. This is made possible by the release of one pivalate oxygen atom from the nickel centre, freeing up an extra site for coordination. While previous studies had involved only replacement of the aquo ligand in the wheels, chelation clearly provides a driving force for this pivalate substitution.

A similar reaction was attempted using a slightly extended version of qpy, bbpe, to form 304. The reaction was carried out in an identical fashion to that used to form 303, but the addition of acetone to the reaction mixture caused no precipitation. A product has been isolated from this reaction by recrystallisation, involving gentle evaporation of an acetone/acetonitrile solution at room temperature. The elemental analyses obtained on
this sample suggested the formation of the desired product was achieved, but a crystal structure is required in order to confirm its structure.

### 3.3.3 Synthesis of Metal Complex Linker Molecules

Various rhenium, ruthenium or iron polypyridyl complexes have been synthesised for use as linker molecules. These were chosen for their expected MLCT photoexcitation and/or redox properties. Some have been prepared following previously published literature procedures, whereas others are new.

![Scheme 14. Synthetic procedure for Re-based linkers.](image)

The synthesis of **305** was achieved following a literature procedure by Thomas et al.\textsuperscript{108} The synthesis of the bbpe analogue, fac-ReCl(CO)\textsubscript{3}(bbpe), **306**, was attempted in an identical fashion (Scheme 14). Characterisation of the product proved very difficult due to poor solubility. Elemental analyses suggest some contamination by the ReCl(CO)\textsubscript{5} starting material, and the solid-state IR spectrum shows two v(CO) bands (one sharp, the other broader) between 2015 and 1840 cm\textsuperscript{-1} as expected for compounds of this type. Repeating this reaction using an excess of bbpe might provide a pure sample of **306**, but as this material could not be further reacted with wheels because of its incredibly poor solubility, this possibility was not further investigated.
The synthesis of \(307\) was achieved using a literature method by Ward et al\(^{107}\) and the product fully characterised (Scheme 15). The same method was applied to form the novel compound \(308\) using bbpe as the incoming ligand in place of qpy. Purification of \(308\) was achieved by column chromatography on silica gel and the product was characterised by \(^1\)H NMR and UV-vis spectroscopies and elemental analyses.

Complex salt \(309\) was synthesised according to a literature method (Scheme 16).\(^{105}\)
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The formation of unsymmetrical Fe\(^{II}\) tris-chelate complexes of type \([\text{Fe}(L)_2(L')_2]^{2+}\) has not been widely reported in the literature. However, two different methods have been used to form such compounds. The first approach\(^{109}\) involves simply reacting a 2:1 ratio of the respective ligands 2,2'-bpy (L) and a variety of extended 2,2'-bpy viologen derivatives (L') with Fe\(^{II}\)SO\(_4\)•7H\(_2\)O in water for 1 h at room temperature to form the desired products by precipitation. The products were obtained in quantitative yields and seemed pure by elemental analysis. However, as the only step of purification was recrystallisation and as there was no evidence of characterisation by \(^1\)H NMR this does not seem like a reliable reference. The second published method\(^{110}\) involves a two-step synthesis at \(-30\) °C to form an intermediate of type \([\text{Fe}^{II}(2,2'-\text{bpy})_2(S)_2]^{2+}\) (where \(S = H_2O\) or MeCN) in situ before addition of the second ligand, an extended 2,2'-bpy derivative, to form the desired product in a yield of approximately 70%.

A simple approach was used to synthesise \(^{311}\), based on the first method reported (Scheme 17). One equivalent of \([\text{Fe}^{II}(H_2O)_6][\text{BF}_4]_2\) was added to a refluxing acetone solution of the two ligands, 2,2'-bpy and bbpe, forming products instantaneously. Purification was achieved in a similar fashion to that used for the Ru\(^{II}\) analogues on silica gel. The separation of similarly coloured bands on the column showed that a mixture of products formed, and the lower than anticipated yield of \(^{311}\) suggests strongly that the previously published work did not in fact produce the expected compounds in pure form simply by recrystallisation. Despite obtaining a good \(^1\)H NMR spectrum of this compound (Section 3.3.5), the elemental analyses obtained are unsatisfactory. Therefore, this product has not been further reacted with wheels.
In a similar fashion, a reaction was attempted to form the qpy analogue, 310. Although a pure product has not yet been isolated, the electrospray mass spectrum of the impure product shows peaks at 822 and 339 corresponding to $[M - PF_6]^+$ and $[M - 2PF_6]^{2+}$, respectively.

### 3.3.4 Synthesis of Systems with Wheels Linked Through Metal Centres

A simple procedure was used to link wheels through 305 (Scheme 18). Purification was achieved by gentle evaporation of an acetone/acetonitrile solution and then a dichloromethane/acetonitrile solution, producing a crystalline material. 312 was characterised using mass spectrometry, elemental analyses, IR and UV/Vis.
spectroscopies and cyclic voltammetry. Unfortunately, no crystal structure has been obtained to date, but the large amount of data obtained on this compound leaves little doubt concerning its structure.

![Scheme 19. Synthesis of Ru-containing linked-wheel systems.](image)

The synthesis of 313 and 314 was achieved simply by reacting a 2:1 ratio of wheel and metal complex at room temperature (Scheme 19). The increased solubilities of the products enabled separation from the starting materials, by first extracting into acetonitrile then into toluene. 313 and 314 were characterised by elemental analyses, mass spectrometry, UV-vis spectroscopy and cyclic voltammetry.

The synthesis of 315 was achieved in a fashion identical to 313 and 314 (Scheme 20), and the product was characterised by elemental analyses, mass spectrometry, cyclic voltammetry and UV-vis spectroscopy.

3.3.5 $^1$H NMR Studies

Figure 59. $^1$H NMR spectrum of 311 in acetone-$d_6$ at 293 K.
1H NMR spectra were recorded for the two novel metal complex salts 308 and 311. The spectrum of 311, shown in Figure 59, displays a series of peaks in the aromatic region, which correlate to the 15 different proton environments of this compound, with a number of signals overlapping.

3.3.6. Mass Spectrometry

Mass spectrometry has been used to identify the linked-wheel compounds. Spectra recorded at The University of Manchester show peaks corresponding to the linking molecule and the purple wheel without a water molecule attached. In collaboration with Dr Lindsay Harding of the University of Huddersfield, the spectra of 302, 303, 312, 313 and 314 have been obtained. As for 219, the molecular ion peak is not observed for 303 as the mass of this product is too high. However, the spectra of the other compounds do show peaks correlating with their molecular masses.

3.3.7. IR Spectroscopy

The IR spectrum of the novel linked-wheel system 312 and can be seen in Figure 60. The major differences in the spectra of that and that of 305 occur above 1600 cm⁻¹, where 312 has extra bands due to the pivalate groups on the wheel. The distinctive $\nu$(C=O) peaks are positioned very similarly in these two compounds, suggesting that the wheels do not exert significant electronic effects on the Re centre.
3.3.8. UV-Visible Spectroscopy

The UV-vis absorption spectra of the various complexes have been recorded in order to observe any d→π* MLCT transitions. These studies have been carried out at 293 K in acetonitrile, except for 305 and 312, which have been studied in dichloromethane due to solubility considerations. The resulting data can be seen in Table 4.

Table 4. UV-vis data for compounds 305, 307–308 and 311–314.

<table>
<thead>
<tr>
<th>Compound</th>
<th>λ_{max}/nm (ε/M^{-1} cm^{-1})</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>305^a</td>
<td>413 (3 900) d→π*</td>
<td></td>
</tr>
<tr>
<td></td>
<td>307 (16 600) π→π*</td>
<td></td>
</tr>
<tr>
<td>307^b</td>
<td>462 (16 500) d→π*</td>
<td>π→π*</td>
</tr>
<tr>
<td></td>
<td>356 (13 000) d→π*</td>
<td>π→π*</td>
</tr>
<tr>
<td></td>
<td>288 (70 300) π→π*</td>
<td>π→π*</td>
</tr>
<tr>
<td></td>
<td>254 (51 100) π→π*</td>
<td>π→π*</td>
</tr>
<tr>
<td>308^b</td>
<td>472 (24 500) d→π*</td>
<td>π→π*</td>
</tr>
<tr>
<td></td>
<td>376 (18 800) d→π*</td>
<td>π→π*</td>
</tr>
<tr>
<td></td>
<td>289 (105 300) π→π*</td>
<td>π→π*</td>
</tr>
<tr>
<td>Compound</td>
<td>Wavelength</td>
<td>Band Type</td>
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<tr>
<td>----------</td>
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</tr>
<tr>
<td>$^{311}$</td>
<td>257 (45 600)</td>
<td>$\pi\rightarrow\pi^*$</td>
</tr>
<tr>
<td>$^{311}$</td>
<td>536 $^c$</td>
<td>d→$\pi^*$</td>
</tr>
<tr>
<td>$^{311}$</td>
<td>410 $^c$</td>
<td>d→$\pi^*$</td>
</tr>
<tr>
<td>$^{311}$</td>
<td>336 $^c$</td>
<td>$\pi\rightarrow\pi^*$</td>
</tr>
<tr>
<td>$^{311}$</td>
<td>300 $^c$</td>
<td>$\pi\rightarrow\pi^*$</td>
</tr>
<tr>
<td>$^{312}$</td>
<td>556 (1 600)</td>
<td>d→d</td>
</tr>
<tr>
<td>$^{312}$</td>
<td>419 (6 600)</td>
<td>d→$\pi^*$</td>
</tr>
<tr>
<td>$^{312}$</td>
<td>308 (25 500)</td>
<td>$\pi\rightarrow\pi^*$</td>
</tr>
<tr>
<td>$^{313}$</td>
<td>563 (2 600)</td>
<td>d→d</td>
</tr>
<tr>
<td>$^{313}$</td>
<td>463 (16 700)</td>
<td>d→$\pi^*$</td>
</tr>
<tr>
<td>$^{313}$</td>
<td>360 (13 100)</td>
<td>d→$\pi^*$</td>
</tr>
<tr>
<td>$^{313}$</td>
<td>288 (71 200)</td>
<td>$\pi\rightarrow\pi^*$</td>
</tr>
<tr>
<td>$^{313}$</td>
<td>250 (72 000)</td>
<td>$\pi\rightarrow\pi^*$</td>
</tr>
<tr>
<td>$^{314}$</td>
<td>555 (2 300)</td>
<td>d→d</td>
</tr>
<tr>
<td>$^{314}$</td>
<td>478 (55 300)</td>
<td>d→$\pi^*$</td>
</tr>
<tr>
<td>$^{314}$</td>
<td>386 (10 100)</td>
<td>d→$\pi^*$</td>
</tr>
<tr>
<td>$^{314}$</td>
<td>289 (12 900)</td>
<td>$\pi\rightarrow\pi^*$</td>
</tr>
</tbody>
</table>

$a$ Solutions in dichloromethane. $^b$ Solutions in acetonitrile. $^c$ Values of $\varepsilon$ have not been calculated due to unsatisfactory elemental analyses.

The data for $^{305}$ agree with those in the literature, with the peak at 307 nm being assigned to a $\pi\rightarrow\pi^*$ transition and that at 413 nm being assigned to a d→$\pi^*$ transition. As can be seen in Figure 61, the position of the MLCT band changes little, but the apparent intensity does increase upon coordination of the wheels. However, the increased $\varepsilon$ value may be attributable to overlap with wheel-based absorption(s). This is an observation that is remarked upon in more detail in Chapter 4. The broad band in the spectrum of $^{312}$ at $\lambda_{\text{max}} \approx 560$ nm arises from d-d transitions within the Cr$_7$Ni wheels. The Cr$_7$Ni wheel precursor shows a band at a very similar wavelength.
The UV-vis spectra of **307**, **308**, **313** and **314** can be seen in Figure 62. In a similar fashion to the previously discussed spectra, addition of wheels to a complex (moving from **307** to **313**) does not significantly alter the position of the MLCT bands, and in this case neither does it appreciably affect the band intensities. A weak band at $\lambda_{\text{max}} \approx 560$ nm in the spectrum of **313** is once again assigned to d-d transitions within the wheels. The spectrum of **308** shows bathochromic shifts in comparison to that of **307**, with changing the ligand from qpy to bbpe also giving a hyperchromic effect. Both of these observations are attributable to the extended $\pi$-conjugation and increased orbital overlap within the ligand.
Figure 62. UV-vis absorption spectra of 307, 308, 313 and 314 in acetonitrile at 293 K.

Figure 63. UV-vis absorption spectrum of 311 in acetonitrile at 293 K.

While valid molar extinction coefficients cannot be derived for 311 due to unsatisfactory elemental analyses, its spectrum (Figure 63) clearly displays two low energy absorptions.
assigned to MLCT transitions. These bands occur at wavelengths that are consistent with data found in the literature for similar [Fe$^{II}$(2,2'-bpy)$_2$(L)]$^{2+}$ complexes (where L is a α-diimine ligand).\textsuperscript{110}

3.3.9. Electrochemistry

Cyclic voltammetric studies have been carried out with the novel linked-wheel systems and their precursors 305, 307–309 and 311–315, and the results are shown in Table 5. The presence of terminal pyridyl groups did create difficulties due to coating of the working electrode, but good data were obtained nonetheless.

<table>
<thead>
<tr>
<th>Compound</th>
<th>$E_{1/2}$/V or $E_{pc}$/V vs. M$^{III/II}$</th>
<th>Ag/AgCl ($\Delta E_g$/mV) $^d$</th>
<th>Ligand Waves $^d$</th>
</tr>
</thead>
<tbody>
<tr>
<td>305 $^a$</td>
<td>$-1.10$</td>
<td>$-1.06$ (80)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>307 $^b$</td>
<td>1.39 (100)</td>
<td>$-1.11$ (60)</td>
<td></td>
</tr>
<tr>
<td>308 $^b$</td>
<td>1.34 (70)</td>
<td>$-1.13$ (60)</td>
<td>$-0.57$</td>
</tr>
<tr>
<td>309 $^a$</td>
<td>1.24 (40)</td>
<td>$-1.04$ (54)</td>
<td>$-1.21$ (76)</td>
</tr>
<tr>
<td>311 $^b$</td>
<td>1.12 (100)</td>
<td>$-0.67$</td>
<td>$-1.11$ (85)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>$-1.39$</td>
</tr>
<tr>
<td>312 $^a$</td>
<td></td>
<td>$-1.06$ (80)</td>
<td></td>
</tr>
<tr>
<td>313 $^b$</td>
<td>1.41 (40)</td>
<td>$-1.09$</td>
<td></td>
</tr>
</tbody>
</table>
The cyclic voltammograms of 305 and its corresponding linked-wheel system 312 can be seen in Figure 64. As anticipated for ReI-diimine systems, no redox activity is observed for the Re centre within the window measured, since such complexes normally show an irreversible ReII/I oxidation above 1.5 V vs. Ag/AgCl. In 305, irreversible processes are observed between −0.5 and −1.2 V, corresponding to ligand-centred reductions, in accord with literature studies. In contrast, 312 shows a single, quasireversible reduction wave with $E_{1/2} = −1.06$ V vs. Ag/AgCl. Therefore, it appears that coordination of the pyridyl N atoms to the purple wheels improves the reversibility of the reductive chemistry of 305.

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<tbody>
<tr>
<td>314</td>
<td></td>
<td></td>
</tr>
<tr>
<td>b</td>
<td>1.35 (70)</td>
<td>−1.13</td>
</tr>
<tr>
<td></td>
<td></td>
<td>−0.87</td>
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<tbody>
<tr>
<td>315</td>
<td></td>
<td></td>
</tr>
<tr>
<td>a</td>
<td>1.30 (90)</td>
<td>−0.80</td>
</tr>
<tr>
<td></td>
<td>−0.95 (80)</td>
<td>−1.17 (70)</td>
</tr>
</tbody>
</table>

*In dichloromethane. b In acetonitrile. c Solutions ca. $10^{-4}$ M in analyte and 0.1 M in [NBu₄][PF₆] at a Pt disc working electrode with a scan rate of 200 mV s⁻¹. Ferrocene reference $E_{1/2} = 0.44$ V, $\Delta E_p = 65$ mV. d $E_{pc}$ value unless stated otherwise.

---

Figure 64. CVs of 305 and 312 in acetonitrile 0.1 M in [NBu₄][PF₆] at 200 mV s⁻¹.
Chapter 3 – Linked Purple Wheel Systems

The cyclic voltammograms of 307 and 313 can be seen in Figure 65. The data obtained for 307 agree with those in the literature. The trace obtained for 313 is of poorer quality when compared with that of 307, probably due to the high molecular mass and lower solubility of 313. However, comparison of the two voltammograms reveals that the Ru$^{III/II}$ $E_{1/2}$ value is not affected significantly by coordination of the wheels, although the ligand-based reduction at $E_{1/2} = -1.11$ V vs. Ag/AgCl for 307 becomes irreversible in 313. This provides an interesting contrast with the behaviour of the Re$^{I}$-containing pair 305 and 312.

![Figure 65. CVs of 307 and 313 in acetonitrile 0.1 M in [NBu$_n$]$_4$[PF$_6$] at 200 mV s$^{-1}$.](image)

The cyclic voltammograms of 308 and 314 show a pattern similar to those obtained for 307 and 313, with a much weaker signal for 314 when compared with 308. Once more the Ru$^{III/II}$ $E_{1/2}$ value is not altered and the ligand based reduction at $E_{1/2} = -1.13$ V vs. Ag/AgCl becomes irreversible upon addition of the wheels. The irreversible reduction wave at $-0.57$ V vs. Ag/AgCl in 308 is attributable to the presence of the ethenylene groups.
Figure 66. CVs of 308 and 314 in acetonitrile 0.1 M in [NBu₄][PF₆] at 200 mV s⁻¹.

Figure 67. CVs of 309 and 315 in dichloromethane 0.1 M in [NBu₄][PF₆] at 200 mV s⁻¹. Inset: Close up of ligand based reductions.

The redox behaviour of 309 and 315 varies once again only in the ligand reduction region (Figure 67). As anticipated from literature results, 309 shows two reversible ligand
based reduction waves. \textbf{315} seems to show two reversible ligand based reductions, at −0.95 V and −1.17 V vs. Ag/AgCl with an irreversible band appearing at -0.80 V.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{Cyclic_voltammogram.png}
\caption{Cyclic voltammogram of 311 in acetonitrile at a scan rate of 200 mV s$^{-1}$.}
\end{figure}

The cyclic voltammogram of 311 (Figure 68) shows a quasi-reversible Fe$^{III/II}$ oxidation wave with $E_{1/2} = 1.12$ V vs. Ag/AgCl. Several ligand-centred reduction processes are also observed, one of which appears to be reversible. As for 308, the irreversible reduction wave at −0.67 V vs. Ag/AgCl in 311 arises from the presence of the ethenylene units.

\subsection*{3.3.10. Luminescence Studies}

Some initial luminescence studies have been run on the Re$^I$ complexes 305 and 312. 305 shows a broad emission band at 650 nm upon excitation at 400 nm, which agrees with literature data and is assigned to emission from a MLCT excited state. In contrast, 312 displays no luminescence upon 400 nm excitation, showing that it is quenched due to the presence of the wheels. Presumably, this observation reflects a shortening of the MLCT excited state lifetime due to nonradiative decay via other electronic excited states, possibly of ligand-field nature. Further, more detailed luminescence studies (also
involving the RuII complexes) will be required in order to clarify the behaviour of the new linked-wheel systems.

3.3.11. X-Ray Crystallography

Crystal structures have been solved for 302 and 303. Both adopt orthorhombic space groups; C2221 for 302 and P22121 for 303.

![Crystal Structure of 302](image)

The crystal structure of 302 can be seen in Figure 69. The two N–Ni bond distances of 2.057(4) Å are similar to those observed in the previously published 4,4′-bpy linked-wheel system. The bpvb linker molecule is not planar and twists about the ethylene bridges with torsion angles of almost 20°.

The structure of 303 (Figure 70) shows an interesting arrangement of wheels not seen previously. The qpy ligand is coordinated to three wheels, with chelation to the central one. All four N–Ni distances are approximately 2.1 Å, showing no significant shortening upon chelation. An equatorial O (*), originally coordinated to the NiII centre, has been
released, leaving a monodentate pivalate group attached to the adjacent Cr$^{III}$ ion. All other bonds in the structure of the chelated wheel remain as expected.

Figure 70. Crystal structure of 303.
3.4. CONCLUSIONS

Three systems linked through purely organic moieties have been synthesised, with 302 and 303 characterised by X-ray crystallography. Two novel transition metal-containing linker molecules (308 and 311) have been prepared using the ligand bbpe, and both have been characterised using $^1$H NMR spectroscopy. 308 and three other Re, Ru or Fe complexes have been reacted with wheels to form linked-wheel systems. Unfortunately, to date no diffraction-quality crystals have been grown of any of the systems containing metal-based linkers, but enough evidence has been gathered to indicate that the desired products have been formed. UV-vis studies show no significant changes in the MLCT absorption spectra due to wheel coordination. However, the emission spectrum of 312 shows quenching of the luminescence observed for the precursor compound 305. Cyclic voltammetry on the linked-wheel systems shows that while the metal-based oxidation processes remain essentially unchanged, in every case the ligand-centred reductions are altered significantly due to the presence of the wheels.

3.5. FURTHER WORK

Ideally, crystal structures are required for the compounds not yet characterised by X-ray diffraction. For the Re-containing linked-wheel system 312, it should be possible to form suitable crystals by subtle alterations to the crystal growing solvents, as the formation of crystalline material has been observed using a dichloromethane/acetonitrile solution. For the Ru/Fe-containing linked-wheel systems, crystal growing may prove to be more challenging due to the relatively high solubilities of these products. Metathesis to a salt, such as a chloride, that would be less soluble in solvents like acetonitrile may give more promising results. Alternatively, using a bulkier anion such as BPh$_4^-$ may facilitate the production of crystals by improving the efficiency of crystal packing.

Studies to assess the switchable properties of the new photo and/or redox-active linked-wheel systems will be a high priority. One way of doing this may be to use an EPR machine in combination with a pulsed laser, in order to cause MLCT excitation at the appropriate wavelength. Time resolved EPR spectroscopy has been used to study short-
lived excited states in the past\textsuperscript{112,113}, with some studies involving the MLCT excitations of Re\textsuperscript{I} and Ru\textsuperscript{II} polypyridyl complexes.\textsuperscript{114}

It will be desirable to complete the synthesis of the series of Fe and Ru polypyridyl systems in order to be able to determine which is the most promising for our purposes. Three compounds in this series have either not yet been synthesised or reacted with wheels (Figure 71). 310 must be isolated to a satisfactorily pure degree, and a method to purify 311 remains to be established. The complex salt [Ru(pytpy)$_2$][PF$_6$]$_2$ is known in the literature.

![Diagram of a complex salt](image)

$\text{Fe}^{2+}$

$\text{Ru}^{2+}$

$n = 0,1$

Figure 71. Remaining Fe/Ru species to be coordinated to purple wheels.
Chapter 4 – Mixed Wheel Systems
4.1 INTRODUCTION

As described in Chapters 2 and 3, developments have been made in the synthesis of linked wheel systems using both mono-substituted green wheels and sugar-templated purple wheels. Until now, however, the two genres of wheels have only been considered separately. Having demonstrated the various capabilities of linking each genre of wheel, it became apparent that purple and green wheels have the potential to be linked together, by coordinating the terminal pyridyl group of the green wheel to the vacant site on the divalent metal in the purple wheel. With the latter being a first row transition metal, it is anticipated that the reaction conditions should not need to be extreme and coordination should occur readily.

This is an interesting series to study, as the length of the link between the two wheels is not the only variable that can be considered. Changing the divalent metal in the Cr₇M wheels may affect the interaction occurring between the wheels, and the coordination of the divalent metal may also play an important role – in the green wheels the link involves both the divalent metal and the adjacent chromium centre, whereas with the purple wheels the link is formed through only the divalent metal.

A series of mixed wheel systems has been synthesised by reacting four different mono-substituted green wheels (203, 204, 206 and 207) with four purple wheels (Cr₇M, where M = Ni, Co, Mn and Zn). A total of fourteen novel systems has been formed, with two of the attempted reactions failing to occur. These linked wheel systems have been characterised by elemental analysis, X-ray crystallography and, in some cases, UV-vis spectroscopy and mass spectrometry. EPR studies on Cr₇Ni-Cr₇Ni linked systems have shown that the strongest interaction occurs when using isonicotinate as a link, with 4-pyridazine carboxylate giving the weakest coupling. Studies have also been undertaken on Cr₇Ni-Cr₇Mn and Cr₇Ni-Cr₇Zn systems, with the former showing interactions and the latter showing none.
4.2 EXPERIMENTAL PROCEDURES

4.2.1 Materials and Procedures

\[\text{[Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})(\text{H}_2\text{O})]\]^{60} and \[\text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPy})]\]^{5}

were synthesised by literature methods. The syntheses of the other green wheels are discussed in Chapter 2 and the Cr$_7$Mn, Cr$_7$Zn and Cr$_7$Co analogues of the purple wheels were synthesised according to unpublished methods by Dr Grigore Timco.

4.2.2 Physical Measurements

Elemental analyses were performed by the Microanalytical Laboratory, University of Manchester. Mass spectrometry was performed either using electrospray on a Micromass Platform spectrometer at The University of Manchester or using a Bruker MicrTOF-Q mass spectrometer with an electrospray ionisation source by Lindsay Harding at the University of Huddersfield. EPR studies were performed by Dr Floriana Tuna and John Machin at The University of Manchester. Q-Band EPR was run on an Elexsys E500 spectrometer equipped with a Bruker 2 tesla magnet. X-ray crystallographic data was collected in Berkeley, California by Dr Simon J. Teat and the structures were solved by Dr Robin Pritchard at The University of Manchester.

4.2.3 Syntheses

\[\text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPy})\text{Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})]\] 401

To a warm solution of \[\text{[Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})(\text{H}_2\text{O})]\] (1.79 g, 0.807 mmol) in acetone (130 ml) was added a warm solution of \[\text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPy})]\] (203) (1.80 g, 0.777 mmol) in acetone (200 ml) and the reaction was heated under reflux for 5 min. The blue-grey solution was then allowed to sit covered at room temperature until the product formed as a blue-grey crystalline material, which was collected by filtration and washed with a small amount of acetone. Measurable crystals were isolated from the reaction mixture prior to filtration. Yield: 1.92 g (55%). Elemental analysis
All of the other compounds were prepared and isolated as blue-grey solids in a manner similar to 401; the quantities of reagents used, yields and characterisation data are listed below.

\[ \text{[Pr}_2\text{NH}_2](\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CCHCHPy})\text{Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})] \]

\[ \text{[Pr}_2\text{NH}_2](\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CCHCHPy})\text{Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})] (204) \text{ (200 mg, 0.085 mmol) in acetone (20 ml); [Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})(\text{H}_2\text{O})] \text{ (207 mg, 0.093 mmol) in acetone (20 ml). Yield: 292 mg (76%).} \]

\[ \text{[Pr}_2\text{NH}_2](\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPyd})\text{Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})] \]

\[ \text{[Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})(\text{H}_2\text{O})] (151 mg, 0.068 mmol) \text{ in acetone (20 ml); [Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPyd})] \text{ (206) (152 mg, 0.066 mmol) in acetone (15 ml). Yield: 154 mg (52%).} \]

\[ \text{[Pr}_2\text{NH}_2][\text{Cr}_7\text{CoF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPy})\text{Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})] \]

\[ \text{[Cr}_7\text{NiF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})(\text{H}_2\text{O})] (157 mg, 0.071 mmol) \text{ in acetone (20 ml); [Pr}_2\text{NH}_2][\text{Cr}_7\text{CoF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPy})] \text{ (207) (151 mg, 0.065 mmol) in acetone (20 ml). Yield: 135 mg (46%).} \]
[Pr₂NH₂][Cr₇NiF₈(O₂CCMe₃)₁₅(O₂CPy)Cr₇CoF₃(O₂CCMe₃)₁₅(EtGlu)] 405

[Cr₇CoF₃(O₂CCMe₃)₁₅(EtGlu)(H₂O)] (155 mg, 0.070 mmol) in acetone (20 ml); 203 (156 mg, 0.067 mmol) in acetone (20 ml). Yield: 256 mg (85%). Elemental analysis calc'd (%) for C₁₇₀H₃₀₄CoCr₁₄F₁₁NiO₆₇: C 45.21, H 6.78, N 0.93, Cr 16.12, Ni 1.30. Found: C 43.25, H 6.51, N 0.94, Cr 15.67, Ni 1.31.

[Pr₂NH₂][Cr₇NiF₈(O₂CCMe₃)₁₅(O₂CCHCHPy)Cr₇CoF₃(O₂CCMe₃)₁₅(EtGlu)] 406

[Cr₇CoF₃(O₂CCMe₃)₁₅(EtGlu)(H₂O)] (150 mg, 0.068 mmol) in acetone (20 ml); 204 (152 mg, 0.065 mmol) in acetone (20 ml). Yield: 197 mg (67%). Elemental analysis calc'd (%) for C₁₇₂H₃₀₆CoCr₁₄F₁₁NiO₆₇: C 45.48, H 6.79, N 0.92, Co 1.30, Cr 16.02, Ni 1.29. Found: C 44.22, H 6.74, N 0.89, Co 1.30, Cr 15.77, Ni 1.22.

[Pr₂NH₂][Cr₇NiF₈(O₂CCMe₃)₁₅(O₂CPyd)Cr₇CoF₃(O₂CCMe₃)₁₅(EtGlu)] 407

[Cr₇CoF₃(O₂CCMe₃)₁₅(EtGlu)(H₂O)] (153 mg, 0.069 mmol) in acetone (20 ml); 206 (153 mg, 0.066 mmol) in acetone (20 ml). Yield: 133 mg (45%). Elemental analysis (%) calc'd for C₁₆₉H₃₀₃CoCr₁₄F₁₁Na₂NiO₆₇: C 44.78, H 6.78, N 1.24, Co 1.31, Cr 16.16, Ni 1.30. Found: C 42.27, H 6.65, N 1.24, Co 1.34, Cr 15.08, Ni 1.14.

[Pr₂NH₂][Cr₇CoF₈(O₂CCMe₃)₁₅(O₂CPy)Cr₇CoF₃(O₂CCMe₃)₁₅(EtGlu)] 408

[Cr₇CoF₃(O₂CCMe₃)₁₅(EtGlu)(H₂O)] (155 mg, 0.070 mmol) in acetone (20 ml); 207 (153 mg, 0.066 mmol) in acetone (20 ml). Yield: 159 mg (53%). Elemental analysis calc'd (%) for C₁₇₀H₃₀₄Co₂Cr₁₄F₁₁N₂O₆₇: C 45.20, H 6.78, N 0.93, Co 2.61, Cr 16.12. Found: C 45.24, H 6.91, N 0.99, Co 2.65, Cr 15.96.

[Pr₂NH₂][Cr₇NiF₈(O₂CCMe₃)₁₅(O₂CPy)Cr₇MnF₃(O₂CCMe₃)₁₅(EtGlu)] 409

[Cr₇MnF₃(O₂CCMe₃)₁₅(EtGlu)(H₂O)] (299 mg, 0.135 mmol) in acetone (20 ml); 203 (151 mg, 0.065 mmol) in acetone (20 ml). Yield: 44 mg (15%). Elemental analysis calc'd
Chapter 4 – Mixed Wheel Systems

(% for C$_{170}$H$_{304}$Cr$_{14}$Mn$_3$Ni$_3$O$_{67}$: C 45.25, H 6.79, N 0.93, Cr 16.13, Ni 1.30. Found: C 45.18, H 6.80, N 0.92, Cr 16.12, Ni 1.38.

$[\text{Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CCHCHPy})\text{Cr}_7\text{MnF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})]$ 410

$[\text{Cr}_7\text{MnF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})(\text{H}_2\text{O})]$ (313 mg, 0.141 mmol) in acetone (15 ml); 204 (153 mg, 0.065 mmol) in acetone (20 ml). Yield: 169 mg (57%). Elemental analysis calcd (%) for C$_{172}$H$_{306}$Cr$_{14}$Mn$_3$Ni$_3$O$_{67}$: C 45.52, H 6.80, N 0.93, Cr 16.04, Ni 1.29. Found: C 44.34, H 6.69, N 0.96, Co 1.31, Cr 16.13, Mn 1.22. Found: C 44.34, H 6.69, N 0.96, Co 1.31, Cr 16.13, Mn 1.14.

$[\text{Pr}_2\text{NH}_2][\text{Cr}_7\text{CoF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPy})\text{Cr}_7\text{MnF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})]$ 411

$[\text{Cr}_7\text{MnF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})(\text{H}_2\text{O})]$ (308 mg, 0.139 mmol) in acetone (20 ml); 207 (151 mg, 0.065 mmol) in acetone (15 ml). Yield: 106 mg (52%). Elemental analysis calcd (%) for C$_{170}$H$_{304}$Co$_{14}$Cr$_{11}$Mn$_3$Ni$_3$O$_{67}$: C 45.24, H 6.79, N 0.93, Co 1.31, Cr 16.13, Mn 1.22. Found: C 45.06, H 6.85, N 0.88, Cr 16.32, Zn 1.44.

$[\text{Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CPy})\text{Cr}_7\text{ZnF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})]$ 412

$[\text{Cr}_7\text{ZnF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})(\text{H}_2\text{O})]$ (306 mg, 0.138 mmol) in acetone (20 ml); 203 (150 mg, 0.065 mmol) in acetone (20 ml). Yield: 112 mg (38%). Elemental analysis calcd (%) for C$_{170}$H$_{304}$Cr$_{14}$F$_{11}$N$_3$NiO$_{67}$Zn: C 45.14, H 6.77, N 0.93, Cr 16.09, Zn 1.45. Found: C 45.06, H 6.85, N 0.88, Cr 16.32, Zn 1.44.

$[\text{Pr}_2\text{NH}_2][\text{Cr}_7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{15}(\text{O}_2\text{CCHCHPy})\text{Cr}_7\text{ZnF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})]$ 413

$[\text{Cr}_7\text{ZnF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})(\text{H}_2\text{O})]$ (0.1567 g, 0.071 mmol) in acetone (15 ml); 204 (0.1520 g, 0.065 mmol) in acetone (20 ml). Yield: 95 mg (32%). Elemental analysis calcd (%) for C$_{172}$H$_{306}$Cr$_{14}$F$_{11}$N$_3$NiO$_{67}$Zn: C 45.41, H 6.78, N 0.92, Cr 16.00, Ni 1.29. Found: C 45.11, H 6.77, N 0.86, Cr 16.05, Ni 1.31.
[Pr₂NH₂][Cr₇Co₇(O₂CCMe₃)₁₅(O₂CPy)Cr₇ZnF₃(O₂CCMe₃)₁₅(EtGlu)] 414

[Cr₇ZnF₃(O₂CCMe₃)₁₅(EtGlu)(H₂O)] (304 mg, 0.137 mmol) in acetone (20 ml); 207 (155 mg, 0.067 mmol) in acetone (20 ml). Yield: 190 mg (63%). Elemental analysis calcd (%) for C₁₇₀H₃₀₄CoCr₁₄F₁₁N₃O₆₇Zn: C 45.14, H 6.77, N 0.93, Cr 16.09, Co 1.30. Found: C 44.99, H 6.86, N 0.93, Cr 16.12, Co 1.37.
4.3 RESULTS AND DISCUSSION

4.3.1 Synthetic Studies

By mixing a one-to-one ratio of reactants in refluxing acetone, the straightforward formation of a mixed wheel system occurs (Scheme 21 and Table 6). Simply leaving the reaction mixture covered for between 1 and 10 days affords diffraction-quality crystals of the desired products.

\[
\text{L} = [\text{Pr}_3\text{NH}_2][\text{Cr}_7\text{MF}_6(\text{O}_2\text{CCMe}_3)_{15}(L)]
\]

When \( M = \text{Ni} \), \( L = \]

\[
\text{L} = [\text{Cr}_7\text{MF}_3(\text{O}_2\text{CCMe}_3)_{15}(\text{EtGlu})]
\]

(\( \text{Cr}_7\text{M} \)) where \( M = \text{Ni}, \text{Co}, \text{Mn} \) or \( \text{Zn} \)

The reaction of 401 was also repeated on a larger scale, where it was found possible to speed up the product formation by stirring the reaction mixture for an hour instead of allowing it to sit covered until crystals formed. The powder formed was proven to be pure by elemental analysis, however crystals suitable for x-ray diffraction were not obtained by this method.
Most of the reactions were successful, but two crystal structures obtained on the products obtained from reactions with the 4-pyd substituted GW 206 showed that the desired mixed-wheel systems did not form. Attempted reactions with the \( \text{Cr}_7 \text{Mn} \) and \( \text{Cr}_7 \text{Zn} \) PWs lead to only co-crystallisation of the starting materials, with no actual links formed between the two wheels.

### 4.3.2 Mass Spectrometry Studies

One method of characterisation used with these wheel systems at The University of Manchester is Electrospray mass spectrometry. As mentioned in Chapters 2 and 3, systems containing two wheels are too large to be detected in their entirety, but it is possible to identify peaks characteristic of various fragments. The spectral region between 2200 and 2400 Da contains peaks relating to \([GW]^+\), \([GWNa]^+\), \([PW - H_2O]^+\) and \([PW - H_2O + Na]^+\). The presence of a peak corresponding to the green wheel does not confirm anything about the linked arrangement of the system, but the presence of a peak for a purple wheel without a water (or acetone) molecule implies an originally linked arrangement.

In order to further characterise this type of linked wheel system by mass spectrometry, a representative example (402) was also sent to Huddersfield to be studied by Dr Lindsay Harding. The results show that as well as the peaks in the 2200–2400 Da region, a large peak is also visible at 4544 Da, indicative of \([\text{MNa}]^+\). This observation confirms that the
wheels remain linked even in solution. It is anticipated that related studies with the other linked-wheel compounds described in this chapter would give similar results.

4.3.3 UV-VIS Spectroscopy

![UV-Vis Spectra](image)

Figure 72. UV-vis spectra of 401 and its components.

The UV-vis spectra of 401 and its component wheels (Figure 72) were run in dichloromethane and directly compared. In all cases, two peaks can be observed between 400 and 700 nm, which are assigned to d-d transitions, and as expected show weak intensities as they are Laporte forbidden. As can be seen in Figure 73, the spectrum for the linked-wheel system corresponds with a pure summation of those of the two starting materials, showing that linking the wheels together has no effect on their electronic transitions.
Figure 73. Additive spectrum of the two starting materials against the spectrum of 401.

The UV-vis spectra of 402 and its starting materials were also studied (Figure 74). Once again an additive effect is observed, so it is anticipated that such behaviour will be seen for all mixed-wheel systems of this type.

Figure 74. UV-vis spectra of 402 and its components.
4.3.4 EPR Studies

While the interactions between these wheels are not strong enough to be detected using SQUID analysis, EPR spectroscopy is a more sensitive technique that can afford useful information.

The EPR spectra of 401–403 can be seen in Figure 75. All three spectra are modelled on the same spin-Hamiltonian parameters (Equation 8), with $S_{\text{eff}} = 1$.

\[
\vec{H}_S = \mu_B \mathbf{B}_g \vec{S} + D \left( \vec{S}_z - \frac{S(S+1)}{3} \right)
\]

As can be seen in Figure 76, the simulated spectrum of 403 does not match well with the experimental spectrum. This observation could result from a number of factors, for example it is possible that the compound is rhombic. An alternative suggestion is that the product was perhaps slightly less crystalline than the other linked-wheel systems. It is known that some compounds give different EPR spectra if they are in a powder as
opposed to a crystalline form, which may allow the parameters to better fit the experimental result.

Figure 76. EPR spectrum of 403 and its simulation using the parameters below.

Table 7. EPR parameters for 401–403.

<table>
<thead>
<tr>
<th></th>
<th>401</th>
<th>402</th>
<th>403</th>
</tr>
</thead>
<tbody>
<tr>
<td>g_x</td>
<td>1.82</td>
<td>1.818</td>
<td>1.824</td>
</tr>
<tr>
<td>g_y</td>
<td>1.82</td>
<td>1.818</td>
<td>1.824</td>
</tr>
<tr>
<td>g_z</td>
<td>1.79</td>
<td>1.8</td>
<td>1.78</td>
</tr>
<tr>
<td>Line Widths (G)</td>
<td>120 (isotropic)</td>
<td>x = y = 100 ≠ z = 120</td>
<td>x = y = 100 ≠ z = 140</td>
</tr>
<tr>
<td>D (cm^{-1})</td>
<td>+0.057</td>
<td>+0.042</td>
<td>+0.036</td>
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</tbody>
</table>

Table 7 shows the parameters used to obtain the three simulated spectra. As explained above, the parameters used to ascertain the simulation for 403 are not very reliable and more work must be undertaken to improve these results. However, from these results it appears that the strongest interaction between wheels is observed for 401, as indicated by its relatively large zero field splitting (D value). The smallest D value, and hence the weakest interaction, is observed for 403; this is interesting as the only structural difference in structure between 401 and 403 is the presence of an N instead of a CH in the ortho position of the pyridyl ring. It may be the case that the greater localisation of electrons within a pyridazine as opposed to a pyridine ring inhibits the interaction.
between the wheels. This hypothesis can be supported by previous structural studies on pyridazine,\textsuperscript{115} which show unexpected changes in bond length around the ring, not seen in a ring with full aromatic delocalisation.

Figure 77 shows the spectra of \textbf{410} and its precursors, together with the summation of the latter spectra. The purple wheel used in this synthesis is Cr\textsubscript{7}Mn, with \(S_{\text{eff}} = 1\) (\(S\) of Mn\textsuperscript{II} = \(5/2\)). The spectrum of \textbf{410} is very different to the sum of those of its starting materials, indicating that some interaction occurs between the wheels. Variable temperature EPR studies will be necessary in order to clarify whether the low-lying \(S_{\text{eff}} = 2\) excited state of the Cr\textsubscript{7}Mn wheel influences the spectra observed.
In contrast, the spectrum of compound 413 (Figure 78) is virtually identical to the sum of those of its starting materials, indicating no interaction occurring between the wheels. The only difference between this and 410 is the divalent metal. Zn\(^{2+}\) has a full d-subshell, as opposed to Mn\(^{2+}\), thus preventing interaction occurring between the two wheels.

4.3.5 X-Ray Crystallography

Crystal structures were obtained for seven of the new linked-wheel systems (401, 402, 405, 407, 409, 412 and 413), confirming their expected molecular structures. Two of these structures (401 and 402) have been finished and are discussed below, while the others are all very similar so were not deemed worthy of further work. For simplification purposes, the \(^1\)Bu groups of the pivalate ligands have not been shown.
Both crystal structures are monoclinic and adopt the $P_{1}2_{1}1$ space group. Both have normal Ni–N bond distances in the range 2.03–2.06 Å. The largest difference between the two structures is the relative arrangement of the wheels. In 401 (Figure 79) the wheels are arranged in a transoid orientation, whereas in 402 (Figure 80) they are positioned cis to each other.
4.4 CONCLUSIONS

Purple and green wheels have been successfully linked together in a simple synthetic procedure, and characterised by elemental analysis, mass spectrometry and, in certain cases, X-ray crystallography. Two reactions involving 206 as a starting material were unsuccessful, maybe as a result of the relatively poor monodentate coordinating abilities of the 4-pyridazine carboxylate ligand.

Interaction between the wheels in these systems is weak, and thus has not been detected using SQUID or INS studies. However interaction has been observed using EPR spectroscopy. The strength of interaction decreases in a predictable manner when the link is extended from ISNA to 4-pyac. Using a 4-pyd ring as a link (in 403) gives the weakest interactions. This observation may be attributable to decreased delocalisation of electrons within the ring, hindering communication between the wheels. Interaction has also been observed via EPR between a Cr7Ni green wheel and a Cr7Mn purple wheel, but no interaction is observed when using a Cr7Zn purple wheel.

4.5 FURTHER WORK

This series of mixed-colour and mixed-metallic wheels merits further studies. To date, studies have involved Cr7M (M = Ni, Co, Mn and Zn) purple wheels, but only Cr7Ni and Cr7Co green wheel systems. By expanding the series of mixed-metal systems to include Cr7Mn and Cr7Zn green wheels it would be possible to assess whether interactions occur when Cr7Zn green wheels are used as opposed to their purple analogues. This may be the case, as in the green wheel systems the linkage is across two metal centres, not only the ZnII ion.

Another possibility would be to synthesise some potentially switchable mixed-wheel systems related to the Ru and Re-containing systems considered in Chapters 2 and 3, some examples of which may be seen in Figure 81. The properties of these species could be compared informatively with those containing two green or two purple wheels.
Figure 81. Some target mixed-wheel systems.
Chapter 5 – Linked Wheels through Hydrogen Bonds
5.1 INTRODUCTION

The first examples of linked wheel systems produced by the Winpenny group involved templating two wheels about a simple linear diamine template, such as 1,8-diaminooctane, 1,9-diaminononane and 1,12-diaminododecane. Reactions were carried out in a fashion identical to the synthesis of individual wheels, allowing the formation of two heterometallic wheels around one template. An example of one of these linked wheel systems can be seen in Figure 82.

Another approach used to link wheels together indirectly was to synthesise a wheel templated about a functionalised secondary amine, such as 4-(ethylaminomethyl)pyridine, and to use the pyridyl group on the template to co-ordinate two wheels to transition metal centres. Primarily, such linked wheel systems were made with first row transition metal dimers, giving products of general formula $M_2(O_2CCMe_3)_4[[PyCH_2NH_2Et][Cr_7NiF_8(O_2CCMe_3)]_{16}]_2$ (where $M = Cu, Ni$ or $Co$). Addition of the pre-made wheels to the metal dimer, $M(O_2CCMe_3)_4(L)_2$ ($L = H_2O$ or $HO_2CCMe_3$) in toluene or diethyl ether, followed by gentle evaporation of the solvents produced X-ray diffraction-quality crystals. An example of such a system can be seen in Figure 83. Further to the synthesis of these compounds, coordination to Ru dimers was also achieved in a similar fashion.
Chapter 5 – Linked Wheels through Hydrogen Bonds

Wheels were also linked through a single Cu centre,\textsuperscript{64} the crystal structure showing a trans coordination geometry (Figure 84). These wheels are clearly too sterically bulky to be linked in a cis conformation when joined by such a small connecting unit.

All of these linked wheel systems were synthesised by using simple procedures and minimal heat, with low boiling solvents. Recrystallisation was the sole means of purification and X-ray diffraction-quality crystals were obtained either directly by gentle
evaporation of the reaction mixture, or as an independent step following removal of the reaction solvent.

A variety of organic compounds and transition metal complexes (Figure 85), which display either redox or photoactivity, were considered as potential linker molecules for the wheels.

![Figure 85. A selection of target linker molecules.](image)

Diquaternary 4,4’-bipyridyl derivatives were considered as potential target linker molecules due to their reversible reductive electrochemistry.\(^8^0\) Such species are best known as viologens as upon undergoing the first of two possible one-electron reductions they become a mono radical cation, which is an intense blue colour. Upon further reduction, a quinoid is produced which is yellow. The first reduction is of particular interest to this project as the production of an unpaired electron may enable an alteration in the communication occurring between the wheels. The other advantage of using viologens as linkers is that their excited state lifetimes are in the nano-second (ns) scale, making them long-lived enough to affect the interaction. There are numerous papers
reporting the synthesis of bis-quaternised bipyridyl derivatives in the literature, making them an ideal starting point for synthetic studies.\textsuperscript{116,117}

Another class of linker template molecules considered are MLCT chromophore-containing Ru\textsuperscript{II} and Re\textsuperscript{I} complexes. It is well established that [Ru\textsuperscript{II}(bpy)\textsubscript{3}]\textsuperscript{2+} and fac-Re\textsuperscript{I}Cl(CO)\textsubscript{3}(bpy) derivatives show long-lived MLCT excited states. [Ru\textsuperscript{II}(bpy)\textsubscript{3}]\textsuperscript{2+} derivatives are also known to undergo fully reversible Ru\textsuperscript{III/II} oxidation processes. Coe \textit{et al.} have synthesised the mono-substituted versions of these target linker molecules using 4-carboxy-4’-methyl-2,2’-bipyridine as a starting material.\textsuperscript{118} There are also procedures in the literature whereby 4,4’-carboxy-2,2’-bipyridine has been used to synthesise a bis-amide system.\textsuperscript{119}

The aims of the project were to identify and synthesise a variety of organic molecules and co-ordination complexes that possess long-lived photo-excited states and/or reversible redox properties. These switchable molecules are intended to act as templates for [Cr\textsubscript{7}NiF\textsubscript{8}(O\textsubscript{2}CCMe\textsubscript{3})\textsubscript{16}] wheels and thus must also contain two terminal amine groups. The molecule may be used as a template to form a linked wheel system directly, or in a substitution reaction to replace the initially used template. A second method to link the wheels together indirectly is to synthesise a wheel templated about an amine with a pendant pyridyl group and to co-ordinate two of these to a metal centre which shows redox and/or photo-activity. Finally, the synthesis and studies of the Cr-Co analogues of these heterometallic wheels is considered, in order to assess the suitability of \textsuperscript{1}H NMR as an analytical tool for these compounds.
5.2 EXPERIMENTAL PROCEDURES

5.2.1 Materials and Procedures

The compounds 4,4’-bis-(carboxy)-2,2’-bipyridine, cis-bis(2,2’-bipyridine)ruthenium (II) dichloride, [4,4’-bis(carboxy)-2,2’-bipyridine]bis(2,2’-bipyridine)ruthenium(II) dichloride, [PyCH₂NH₂Et][Cr₇NiF₈(O₂CCMe₃)₁₆], [Pr₂NH₂][Cr₇NiF₈(O₂CCMe₃)₁₆], [PyCH₂NH₂Et][Cr₇NiF₈(O₂CCMe₃)₁₆(O₂CPy)]², [PyCH₂NH₂Et][Cr₇CoF₈(O₂CCMe₃)₁₆]⁶, fac-ReCl(CO)₃(bpy)¹²¹ and fac-[Re(MeCN)(CO)₃(bpy)] [CF₃SO₃]¹²² were synthesised by previously established methods. They were obtained in yields similar to those of the literature preparations and their identities were confirmed by ¹H NMR spectroscopy, electrospray mass spectrometry and elemental analysis. All other starting materials were purchased commercially and used as supplied.

5.2.2 Physical Measurements

All ¹H NMR spectra were recorded on Bruker Ultrashield 500 MHz or 300 MHz spectrometers. Paramagnetic ¹H NMR spectra were run by Dr Louise Natrajan and Mr Thomas Faust on a Bruker Advance 400 MHz spectrometer at The University of Manchester. Mass spectra were obtained using either electrospray or MALDI analysis on a Micromass Platform spectrometer. Elemental analyses were performed by the Microanalytical Laboratory, University of Manchester. IR data were recorded using a Varian Bio-Rad Excalibur series IR instrument and UV data was recorded on a Shimadzu UV-2401 PC UV-VIS spectrophotometer. Cyclic voltammetry was run in 0.1 M [NBu₄][PF₆] in acetonitrile at a Pt disc working electrode with a scan rate of 200 mV s⁻¹.
5.2.3 Syntheses

N,N’-Bis-(2-aminoethyl)-4,4’-bipyridinium tetrabromide 501

4,4’-Bipyridine (255 mg, 1.63 mmol) was added to a refluxing solution of 2-bromoethylamine hydrobromide (926 mg, 4.52 mmol) in acetonitrile (25 ml) and the reaction refluxed for 96 h, affording a yellow precipitate. The solution was allowed to cool to room temperature and then filtered. The precipitate was washed with acetonitrile and diethyl ether and then allowed to dry under vacuum. Purification was achieved by dissolving the impure product in a 2:1 mixture of water and DMF at 80 °C and then adding acetone to give the pure product as a precipitate, which was once again collected by filtration, washed with acetone and dried under vacuum. Yield: 517 mg (56%). $^1$H NMR (D$_2$O) δH/ppm: 3.68 (4H, t, $J = 4.0$ Hz, CH$_2$), 5.00 (4H, t, $J = 8.0$ Hz, CH$_2$), 8.55 (4H, d, $J = 6.9$ Hz, bpy), 9.14 (4H, d, $J = 6.4$ Hz, bpy). Elemental analysis calcd. (%) for C$_{14}$H$_{22}$Br$_4$N$_4$•H$_2$O: C 28.79, H 4.14, N 9.59. Found: C 28.58, H 3.98, N 9.45.

N-(2-aminoethyl)-4,4’-bipyridinium dibromide 502

4,4’-Bipyridine (817 mg, 5.23 mmol) was added to a refluxing solution of 2-bromoethylamine hydrobromide (372 mg, 1.82 mmol) in acetonitrile (25 ml) and the solution was refluxed for 18 h, affording a cream coloured precipitate. The solution was allowed to cool to room temperature and then filtered. The precipitate was washed with acetonitrile and diethyl ether and then allowed to dry under vacuum. Purification was achieved by dissolving the impure product in minimal water and then adding acetone to give a precipitate, which was collected by filtration, washed with acetone and dried under vacuum. Yield: 515 mg (75%). $^1$H NMR (D$_2$O) δH/ppm: 3.65 (2H, t, $J = 7.1$ Hz, CH$_2$), 4.93 (2H, t, $J = 6.8$ Hz, CH$_2$), 7.83 (2H, d, $J = 6.5$ Hz, bpy), 8.40 (2H, d, $J = 7.1$ Hz, bpy), 8.69 (2H, d, $J = 5.8$ Hz, bpy), 8.96 (2H, d, $J = 7.1$ Hz, bpy). +ve electrospray: $m/z$: 200.2 [M – 2Br]$^{2+}$. –ve electrospray: $m/z$: 279.1 [M – Br$^-$], 360.3 [M – H]. Elemental analysis calcd. (%) for C$_{12}$H$_{15}$Br$_2$N$_3$•H$_2$O: C 38.02, H 4.52, N 11.08. Found: C 37.85, H 4.12, N 11.08.
Chapter 5 – Linked Wheels through Hydrogen Bonds

N-(2-aminoethyl)-trans-1,2-bis(4-pyridyl)ethylene dibromide 503

This compound was prepared and purified in a manner identical to N-(2-aminoethyl)-4,4'-bipyridinium dibromide, but using E-1,2-bis(4-pyridyl)ethylene (632 mg, 4.20 mmol) instead of 4,4'-bipyridine and 2-bromoethylamine hydrobromide (293 mg, 1.43 mmol) in acetonitrile (35 ml). A cream coloured solid was obtained. Yield: 166 mg (29%); \(^1\)H NMR (D\(_2\)O) \(\delta_{\text{H}}/\text{ppm}: 3.60 (2\text{H, t, } J = 6.7\ \text{Hz, CH}_2), 4.82 (2\text{H, t, } J = 7.1\ \text{Hz, CH}_2), 7.47 (1\text{H, d, } J = 16.3\ \text{Hz, CH}), 7.59 (2\text{H, d, } J = 3.6\ \text{Hz, bpy}), 7.66 (1\text{H, d, } J = 16.5\ \text{Hz, CH}), 8.10 (2\text{H, d, } J = 6.2\ \text{Hz, bpy}), 8.45 (2\text{H, d, } J = 5.3\ \text{Hz, bpy}), 8.73 (2\text{H, d, } J = 8.9\ \text{Hz, bpy}).\)

Elemental analysis calcd. (%) for C\(_{14}\)H\(_{17}\)Br\(_2\)N\(_3\)•0.5H\(_2\)O: C 42.45, H 4.58, N 10.61. Found: C 42.42, H 4.21, N 10.60.

2-(tert-But oxy carbonylamino)ethylammonium)4,4'-bis-(carboxylato)-2,2'-bipyridine 505

Thionyl chloride (2 ml) was added dropwise over 10 min to a suspension of 4,4'-bis-(carboxy)-2,2'-bipyridine (122 mg, 0.500 mmol) in dichloromethane (2 ml). The mixture was stirred at room temperature for 5 h and the solvent was removed under vacuum. The crude acid chloride was dissolved in dry dichloromethane (10 ml) and added over approximately 1 min to a solution of N-(tert-butoxycarbonyl)-1,2-diaminoethane (243 mg, 1.52 mmol) and triethylamine (327 mg, 3.23 mmol) in dry dichloromethane (10 ml). The reaction was allowed to stir for 4 h, during which time the product formed as a pink precipitate. The precipitate was collected by filtration, washed with dichloromethane and diethyl ether and dried under vacuum. Yield 168 mg (60%); \(^1\)H NMR ((CD\(_3\))\(_2\)SO) \(\delta_{\text{H}}/\text{ppm: 1.36 (18H, s, } ^1\text{Bu}), 2.85 (4\text{H, t, } J = 5.5\ \text{Hz, CH}_2), 3.18 (4\text{H, d, } J = 4.8\ \text{Hz, } \text{CH}_2), 7.14 (2\text{H, br s, NH}), 7.73 (2\text{H, d, } J = 3.8\ \text{Hz, bpy}), 8.65 (2\text{H, d, } J = 7.6\ \text{Hz, bpy}), 8.76 (2\text{H, s, bpy}).\)

\(+\text{ve electrospray: } m/z = 161\) (protonated N-(BOC)-1,2-diaminoethane). \(-\text{ve electrospray: } m/z = 243\) (mono-deprotonated 4,4'-bis-(carboxy)-2,2'-bipyridine). \(\nu_{\text{max}}/\text{cm}^{-1}: 3236\ (\text{w}, \nu_{\text{OH}}), 3147\ (\text{m}, \nu_{\text{CH}}), 3068\ (\text{m}, \nu_{\text{CH}}), 2994\ (\text{m}, \nu_{\text{CH}}), 2881\ (\text{m}, \nu_{\text{CH}}), 2136\ (\text{w}, \nu_{\text{C=O}}), 1685\ (\text{s}, \nu_{\text{C=O}}), 1643\ (\text{s}, \nu_{\text{C=O}}), 1582\ (\text{s}, \nu_{\text{C=O}}), 1542\ (\text{s}).\)

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4,4′-bis-(carbonylamino(2-(tert-butoxycarbonylamino)ethyl))-2,2′-bipyridyl 506

N-(tert-butoxycarbonyl)-1,2-diaminoethane (397 mg, 2.48 mmol), 1-hydroxybenzotriazole (HOBT, 452 mg, 3.35 mmol) and N,N′-disopropylcarbodiimide (DIC, 1 ml, 6.62 mmol) were added to a stirred solution of 4,4′-bis(carboxy)-2,2′-bipyridine (256 mg, 1.05 mmol) in dry DMSO (12.5 ml). The reaction was allowed to stir for 24 h, during which time the product formed as a precipitate. The precipitate was collected by filtration, washed with DMSO then acetone, affording the product as a white powder which was dried under vacuum. Yield 354 mg (60%); 1H NMR ((CD3)2NC(D)O) δH/ppm: 1.58 (18H, s, t-Bu), 3.53 (4H, m, CH2), 3.73 (4H, m, CH2), 7.15 (2H, m, NH), 8.12 (2H, m, bpy), 9.08 (2H, d, J = 5.5 Hz, bpy), 9.13 (2H, s, bpy), 9.25 (2H, m, NH). MALDI-MS: m/z = 529 [MH]+, 551 [MNa]+, 656, 445, 413, 379, 301, 250. νmax (cm−1): 3355 (s), 3303 (s), 3061 (w), 2983 (m), 2936 (m), 1695 (s) (C=O), 1648 (s) (C=O), 1592 (m), 1537 (br, s). Elemental analysis calcd (%): C 57.50, H 6.91, N 15.01. Found: C 57.45, H 6.89, N 15.34.

(4,4′-Bis(carboxyl(2-(tert-butoxycarbonylamino)ethyl))-2,2′-bipyridine)bis(2,2′-bipyridine)ruthenium(II) bis(hexafluorophosphate) 509

4,4′-Bis-(carbonylamino(2-(tert-butoxycarbonylamino)ethyl))-2,2′-bipyridyl•0.4Me2SO (403 mg, 0.775 mmol) was added to cis-RuIICl2(bpy)•2H2O (454 mg, 0.812 mmol) in 9:1 ethanol/water (70 ml). The resulting solution was heated under reflux overnight to give a bright red solution. The solvents were removed under vacuum to give the product as the slightly impure chloride salt. Purification was effected by precipitation from minimal water with an excess of aqueous NH4PF6. The solid was then filtered off and washed with water to give the product as an orange red powder. Yield: 757 mg (74%); 1H NMR ((CD3)2CO) δH/ppm: 1.35 (18H, s, t-Bu), 3.32 (4H, d, J = 5.3 Hz, CH2), 3.53 (4H, d, J = 6.6 Hz, CH2), 6.24 (2H, m, NH), 7.60 (4H, m, bpy), 7.94 (2H, d, J = 4.5 Hz, bpy), 8.09 (4H, dd, J1 = 5.9 Hz, J2 = 4.9 Hz), 8.27 (6H, m, bpy) 8.44 (2H, s, NH), 8.85 (4H, d, J = 8.1 Hz, bpy), 9.18 (2H, s, bpy). +ve electrospray: m/z = 1087 [(M – PF6)]2+, 470 [(M–2PF6)]3+. –ve electrospray: m/z = 145 [PF6]-. νmax (cm−1): 3435, 2978, 1670 (C=O), 839 (PF6). λmax (ε/mol⁻¹ dm³ cm⁻¹): 466 (14 100), 351 (9 800), 288 (66 500), 247 (31 400).
Elemental analysis calcd. (%) for C_{46}H_{52}F_{12}N_{10}O_{6}P_{2}Ru•H_{2}O: C 44.20, H 4.35, N 11.21.  
Found: C 43.91, H 4.16, N 11.08.

Tricarbonyl(4,4'-Bis(carboxyl(2-(tert-butoxycarbonylamino)ethyl)-2,2'-bipyridine)chlororhenium(I) 511

4,4'-Bis-(carbonylamino(2-(tert-butoxycarbonylamino)ethyl)-2,2'-bipyridyl•0.4Me_{2}SO  
(184 mg, 0.329 mmol) was added to Re{Cl(CO)}_{5} (105 mg, 0.290 mmol) in methanol (15 ml) and the solution heated under reflux for 6 h, during which time it became a bright orange colour.  
The reaction was allowed to cool and the solvent was removed under vacuum.  
Purification was achieved by reprecipitation from DMF and water, allowing the slightly impure product to be collected as a bright orange solid, washed with water and then dried under vacuum.  
Yield: 206 mg (85%);  
{^{1}H} NMR ((CD_{3})_{2}NCDO) \delta_{H}/ppm: 1.58 (18H, s, \text{tBu}), 3.54 (4H, d, J = 7.5 Hz, CH_{2}), 3.75 (4H, d, J = 6.2 Hz, CH_{2}), 7.18 (2H, s, broad NH), 8.43 (2H, d, J = 6.2 Hz, bpy), 9.40 (2H, s, bpy), 9.49 (2H, s, NH), 9.54 (2H, d, J = 5.0 Hz, bpy).  
+ve electrospray: \text{m/z} = 857 [\text{MNa}^+].  
–ve electrospray: \text{m/z} = 833 [\text{M} – \text{H}].  
\nu_{\text{max}} (\text{cm}^{-1}): 2026 (\text{CO}), 1919 (\text{CO}), 1880 (\text{CO}), 1656 (\text{CO}).  
\lambda_{\text{max}} (\text{e/mol}^{\text{-1}} \text{dm}^{\text{3}} \text{cm}^{-1}): 398 (4 400), 304 (16 800), 245 (26 400).  
Elemental analysis calcd. (%) for C_{29}H_{36}ClN_{6}O_{9}Re•3H_{2}O: C 39.21, H 4.77, N 9.46, Re 20.96.  
Found: C 39.19, H 4.55, N 9.34, Re 20.89.

Reaction of Re{Cl(CO)}_{5} and [PyCH_{2}NH_{2}Et][Cr_{7}NiF_{8}(Me_{3}CCO)_{16}] 516

Re{Cl(CO)}_{5} (37.9 mg, 0.105 mmol) and [PyCH_{2}NH_{2}Et][Cr_{7}NiF_{8}(Me_{3}CCO)_{16}] (508 mg, 0.218 mmol) were refluxed in toluene (10 ml) for 32 h before the solvent was removed under vacuum.  
The impure product was placed on an activated neutral alumina column and eluted using toluene.  
The product was the first green band to be removed, leaving the green starting material on the column.  
Yield: 126 mg.
Chapter 5 – Linked Wheels through Hydrogen Bonds

\[ \text{Cu}^{II}_2(\text{O}_2\text{CCMe}_3)_4(\text{[PyCH}_2\text{NH}_2\text{Et}][\text{Cr}_7\text{CoF}_8(\text{Me}_3\text{CCO}_2)_16])_2 \]

\[ \text{Cr}^{III}_3\text{F}_3\cdot4\text{H}_2\text{O} \ (3.02 \text{ g, 16.7 mmol}), \text{4-(ethylaminomethyl)pyridine} \ (0.336 \text{ g, 2.47 mmol}) \]

and pivalic acid (21.6 g, 213 mmol) were heated at 140 °C for 5 h. \([\text{Co}^{II}_7(\text{Me}_3\text{CCO}_2)_2]_n\) (0.660 g) was then added and the reaction was further heated for 24 h at 160 °C. The solution was cooled to room temperature and acetonitrile (100 ml) was added. After stirring for 2 h, the green precipitate was collected by filtration and washed with acetonitrile, then dried in a desiccator overnight. The solid was added to hexane (80 ml) and the mixture filtered to remove undissolved material. To this solution was added \(\text{Cu}^{II}_2(\text{O}_2\text{CCMe}_3)_4(\text{HO}_2\text{CCMe}_3)_2\) (0.526 g, 0.71 mmol) dissolved in a minimum amount of hexane. The reaction mixture was stirred overnight at room temperature and then allowed to stand for a further 24 h, affording a green precipitate. The product was collected by filtration, washed with hexane until the filtrate became colourless and then dried under vacuum. Yield: 2.13 g (64% wrt Cu starting material). Elemental analysis calcd. (%) for \(\text{C}_{196}\text{H}_{350}\text{Co}_2\text{Cu}_2\text{Cr}_{14}\text{F}_{16}\text{N}_4\text{O}_{72}\): C 45.34, H 6.79, N 1.08, Cr 14.02. Found: C 45.31, H 7.02, N 1.07, Cr 13.60.

\[ \text{[PyCH}_2\text{NH}_2\text{Et}][\text{Cr}_7\text{CoF}_8(\text{Me}_3\text{CCO}_2)_16]\]

To \(\text{Cu}^{II}_2(\text{O}_2\text{CCMe}_3)_4(\text{[PyCH}_2\text{NH}_2\text{Et}][\text{Cr}_7\text{CoF}_8(\text{Me}_3\text{CCO}_2)_16])_2\) (2.00 g, 0.38 mmol) dissolved in thf (60 ml) was added triethylamine (5 drops) and acetylacetone (4 ml). The mixture was heated at 60 °C for 15 min and then stirred at room temperature for a further 24 h. The solvent was removed under vacuum and the impure reaction mixture was washed with acetonitrile and filtered until the filtrate became colourless. The precipitate was then dried overnight under vacuum and recrystallisation was effected by evaporation of a toluene:thf (2:1) mixture at room temperature. Yield: 1.71 g (97%). +ve electrospray: \(m/z: 2352.6 [\text{MNa}]^+\). –ve electrospray: \(m/z: 2329.7 [\text{M – H}]^-, 2191.5 [\text{M – PyCH}_2\text{NH}_2\text{Et}]^-.\) Elemental analysis calcd. (%) for \(\text{C}_{88}\text{H}_{157}\text{CoCr}_7\text{F}_8\text{N}_2\text{O}_{32}\): C 45.36, H 6.79, N 1.20. Found: C 45.11, H 6.88, N 1.12.
N-(3-carboxyethyl)-4,4’-bipyridine hexafluorophosphate 519

To a refluxing solution of 3-bromopropionic acid (275 mg, 1.60 mmol) in acetonitrile (25 ml) was added 4,4'-bipyridine (769 mg, 4.80 mmol) in acetonitrile (5 ml). The solution was refluxed for 18 h affording a pale yellow precipitate, which was collected by filtration and washed with acetonitrile and diethyl ether before being dried under vacuum. Yield: 374 mg (76%). $^1$H NMR ((CD$_3$)$_2$CO) $\delta$/ppm: 3.23 (2H, t, $J$ = 6.2 Hz, CH$_2$), 5.02 (2H, t, $J$ = 6.2 Hz, CH$_2$), 7.87 (2H, d, $J$ = 6.0 Hz, bpy), 8.55 (2H, d, $J$ = 6.6 Hz, bpy), 8.75 (2H, d, $J$ = 6.0 Hz, bpy), 9.24 (2H, d, $J$ = 6.6 Hz, bpy). Elemental analysis calcd (%) for C$_{13}$H$_{13}$BrN$_2$O$_2$•H$_2$O: C 39.81, H 3.19, N 7.14. Found: C 40.16, H 3.28, N 7.43.
5.3 RESULTS AND DISCUSSION

5.3.1 Synthetic Studies

5.3.1.1 Synthesis of Linker Molecules

The first series of linker molecules considered are diquaternised derivatives of 4,4’-bipyridine (4,4’-bpy) or E-1,2-bis-(4-pyridyl)ethylene (bpe). Such molecules may show two reversible one-electron reduction processes. Related monoquaternised derivatives could also prove useful for our purposes, because it should be possible for their nonquaternised end to be complexed to redox and/or photoactive metal centres in order to link two wheels. Scheme 22 shows the syntheses of both di- and monoquaternised derivatives.

Scheme 22. Reactions used to synthesise quaternised 4,4’-bpy and bpe derivatives.

A previously published paper claimed that N,N’-bis-(2-aminoethyl)-4,4’-bipyridyl dibromide had been synthesised by reacting 4,4’-bpy with 2-bromoaminoethane hydrobromide in 50:50 DMF/water. This method was attempted, using 2.5 equivalents of 2-bromoaminoethane hydrobromide to 1 equivalent of 4,4’-bpy, but results were unsatisfactory. Upon refluxing the reaction for a period of two days, a colour change was observed. The solution became a dark blue colour, indicating that the desired product may have formed (when such diquaternised bpy derivatives undergo a one-electron
reduction to a mono-cation radical they become a distinctive intense blue). Upon cooling, the solution reverted back to a yellow colour suggesting the reduction was reversible. However, after working up the reaction it appeared that a variety of products was formed as the \(^1\)H NMR spectrum showed several sets of peaks. Furthermore, the impure reaction yield was low so the pure product yield would have been unsatisfactory.

As a result, steps were taken to improve on the proposed synthesis adapting from another literature procedure to prepare viologens,\(^{124}\) this time using acetonitrile as the solvent. The reaction was attempted over a variety of times, from 8 to 96 h, and also using added AgBF\(_4\), the formation of highly insoluble AgBr helping to drive the reaction. The use of AgBF\(_4\) did appear to increase the speed of the reaction, but it then became very difficult to remove all traces of AgBr. By heating the reaction without AgBF\(_4\) for 96 h, the diquaternised product \(\text{501}\) was obtained in a 56% yield. Comparing the results of other reactions attempted over shorter periods of time indicates that the monoquaternised product is formed initially and precipitates out of solution, but its partial solubility in acetonitrile allows the eventual formation of the diquaternised product over longer reaction times. An identical reaction was attempted using bpe in place of 4,4′-bpy, however the diquaternised product was not formed irrespective of reaction timescales. Purification of \(\text{501}\) was achieved by dissolution of the product in a warm mixture of water and DMF and precipitation with acetone, leaving the monoquaternised product in solution.

The synthesis of the monoquaternised compound was achieved by altering the ratio of reactants from that described above to 1 equivalent of 2-bromoaminoethane hydrobromide and 3 equivalents of 4,4′-bpy. Further to the discovery in the previous synthesis that the monoquaternised product \(\text{502}\) is poorly soluble in acetonitrile, the reaction was heated for only 18 h, during which time the product formed as a pale cream coloured precipitate in a 78% yield. A similar reaction using bpe in place of 4,4′-bpy gave the product \(\text{503}\) in a yield of 30%. In both cases purification was achieved by a water/acetone reprecipitation.
Attempts to crystallise the dication 502 produced some interesting results. Initial attempts to crystallise the bromide salt of this product were not successful so steps were taken to metathesise to the PF$_6^-$ and BPh$_4^-$ salts. In both cases, upon addition of the relevant aqueous salts the expected precipitation did not occur. Consequently, extractions into ethyl acetate were carried out. Diffraction-quality crystals were obtained from both salts, by vapour diffusion of diethyl ether into acetone solutions. However, the structures revealed that the terminal amine groups had reacted with the acetone to form Schiff bases (Scheme 23).

4,4'-Bis-(carbonylaminoethylamine)-2,2'-bipyridyl was viewed as an attractive synthetic target as it has two terminal amine groups and therefore may act as a suitable template molecule for connecting two heteropolymetallic rings. These rings may be attached either to the free pro-ligand or to metal complexes containing this ligand, with facile and reversible redox chemistry of the linkage potentially allowing switching of the magnetic coupling between the rings. Scheme 24 shows the routes attempted to prepare 507.
4,4'-Bis-(carboxy)-2,2'-bipyridine 504 was synthesised via a literature procedure.\textsuperscript{120} Further to a previous report\textsuperscript{118}, a two-step synthesis to form 506 was attempted. Step one involved reacting 504 with thionyl chloride at room temperature to generate the carbonyl chloride intermediate which was then reacted in step two with N-BOC-ethylenediamine to form the product. However, the product obtained was the salt 505. This product, while not of direct relevance to the project has been fully characterised and X-ray diffraction-quality crystals were obtained by vapour diffusion of diethyl ether into a DMSO solution.

A potential reason for the formation of 505 as opposed to 506 is that the first step of the reaction had not gone to completion. On reviewing the literature,\textsuperscript{125,126} it has been claimed that the diacid 504 requires heating under reflux in thionyl chloride for at least 4 hours to achieve the successful formation of the acid chloride intermediate.
The second method attempted was adapted from a literature procedure\textsuperscript{127} and involved the use of the popular amide-coupling reagents diisopropylcarbodiimide (DIC) and 1-hydroxybenzotriazole (HOBT). The C=N bonds of the carbodiimide come under nucleophilic attack from the carboxylic acid group(s), and the reaction is driven by the formation of a highly stable urea by-product. The mechanism of this reaction is depicted in Scheme 25.

![Scheme 25. Mechanism of amide bond formation.](image)

This reaction was carried out in DMF initially, resulting in a precipitate which was shown by \textsuperscript{1}H NMR analysis to be a mixture of 505 and a monosubstituted BOC product (Figure 86). The reaction solvent was then changed to DMSO and upon stirring overnight at room temperature the desired product 506 formed in a 64% isolated yield. This compound has been fully characterised.

![Figure 86. Mono-substituted product formed as a by-product of DMF reaction.](image)
The terminal amine groups of 506 were deprotected by stirring for two hours at room temperature in a mixture of dichloromethane and trifluoroacetic acid. The $^1$H NMR spectrum of the product clearly shows that the 'Bu groups have been removed, but no new signals are observed for the -NH$_2$ groups. Problems have also been encountered with the hygroscopic nature of the deprotected compound. One reason for this is the likely formation of the TFA salt of the protonated amine instead of the neutral amine, which may occur due to the large excess of trifluoroacetic acid used in the deprotection step. As a result, no further characterisation has been achieved for this compound (507).

Scheme 26 shows the two approaches used to synthesise 509. The precursors cis-Ru$^{II}$ (bpy)$_2$Cl$_2$·2H$_2$O and 508 were prepared according to literature methods. The first route used to form the complex 509, which included the direct addition of N-BOC-ethylenediamine to 508, was explored due to difficulties encountered initially in synthesising the compound 506.

The BOC protected complex 509 was formed using a method similar to that used to synthesise 506. This reaction was carried out in DMF and the product was isolated as its PF$_6$ salt. Purification was carried out by eluting with an acetonitrile/water/KNO$_3$ mixture on a silica gel column and the product was obtained in a 45% yield. The $^1$H NMR and mass spectra are in agreement with the proposed structure. The elemental analysis however indicates that the product is still not fully pure, even when accounting for one equivalent of KPF$_6$. 

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Upon completion of the synthesis of the pro-ligand, 506, it was decided to try reacting this compound directly with cis-Ru(bpy)$_2$Cl$_2$·2H$_2$O following a method similar to that used by Beer et al.$^{119}$ Purification of the product was achieved by anion metathesis, giving a 48% isolated yield. The $^1$H NMR spectrum shows this material to be as pure as that obtained from the first attempt using column chromatographic purification. In
addition, elemental analysis shows the product to be pure and this method involving the pre-formed protected pro-ligand appears to be superior to the first approach, as the requirement for column chromatography is eliminated.

Once again, treatment of the BOC-protected complex with TFA in dichloromethane produced an extremely hygroscopic product (510). The $^1$H NMR spectrum of this product shows that it is impure, which could be as a result of a mixture of charged and non-charged species being formed from the TFA treatment.

The synthesis of 511 (Scheme 27) was achieved in a fashion similar to a reaction reported previously by Coe et al.$^{118}$ Re$^I$Cl(CO)$_5$ was reacted with 506 in a 1:1 ratio by heating under reflux in methanol for 5 hours. After reprecipitation from DMF and water, the $^1$H NMR spectrum implied the formation of a pure product, although the elemental analysis suggested the presence of remaining Re$^I$Cl(CO)$_5$ starting material. Hence, the reaction was repeated using a slight excess of the pro-ligand, affording a pure sample of the desired product 511.

### 5.3.1.2 Attempted formation of linked wheel systems

The first approach considered when attempting to link Cr$_7$Ni wheels together was to substitute the template of a pre-formed ring for one of the target linkers. The only such template-substitution reaction attempted previously had involved using the incoming template as a solvent and thus in a many-fold excess. It was decided to investigate whether it may be possible to tailor this reaction to allow substitution of templates when
using smaller quantities of replacing units, perhaps even less than one equivalent. The approach considered is shown in Scheme 28.

Scheme 28. Attempted substitution of templates to form a linked wheel system.

The starting wheel compound 512 was synthesised in a fashion similar to other wheels by 2-phenethylamine as a template. This starting material was chosen for two reasons. Firstly, its synthesis is straightforward, requiring no column chromatographic purification, and gives a high yield. Second, the use of a primary amine as opposed to a secondary amine as a template should facilitate the swapping for another molecule. The idea for this procedure was to treat 512 with a base, [Bu₄N]OH, for an hour to form the intermediate 513 by deprotonating the amine about which the wheel is templated, causing the wheel to “slip off” the template and become associated with Bu₄N⁺ instead. The wheel should not be able to sit around the very bulky Bu₄N⁺ cation and therefore, upon addition of the new linking molecule, the wheel should associate itself instead with the target template via hydrogen bonding to the protonated amine.

Mass spectrometry studies suggest that the first step of the reaction, the removal of the wheel from the template and subsequent association with Bu₄N⁺, was successful. However, following the second step by TLC suggested that only a minimal amount of the desired product was in fact formed. It was decided at this point that the method was too inefficient to warrant further study as potentially more fruitful avenues of investigation were becoming available. It appears that a reasonable rate of substitution requires the linker molecule to be present in a large excess and therefore this method would not be cost effective for more expensive metal complex linkers.

The next approach to the formation of linked wheel systems was to attempt the synthesis of wheels directly about the “linker molecules” in a fashion used for simple linear
diamines previously. The first compound considered in this regard was the monoquaternised product 502, as shown in Scheme 29.

Scheme 29. Direct synthesis of a linked wheel system.

A reaction was attempted by adapting previous syntheses. $\text{Cr}^{III} \cdot 4\text{H}_2\text{O}$, $2\text{Ni}^{II} \cdot \text{CO}_3 \cdot 3\text{Ni}^{II}(\text{OH})_2 \cdot 4\text{H}_2\text{O}$ and 502 were heated in pivalic acid at 160 °C for 24 h, then cooled before acetonitrile was added to give a precipitate. The resultant mixture was found to contain simply the starting compounds, probably due to a lack of dissolution of 502 in the pivalic acid. With no template, the wheel cannot form. One potential reason for this failure is that all previously used templates are neutral prior to protonation of the amine group, whereas these viologen-type molecules are already doubly charged and thus their solubility properties may be quite different.

The next molecule to be used as a potential template is the pro-ligand 506. Problems had been experienced when attempting to deprotect the terminal amine groups of this compound in TFA, but this deprotection step is acid-catalysed so it was decided to attempt this in situ as the reaction solvent for wheel formation is an acid (Scheme 30). Once again, this reaction returned only starting materials as the BOC-protected ligand is insoluble in pivalic acid, meaning that pivalic acid is incapable of deprotecting the ligand.

Scheme 30. Attempted synthesis of linked wheel system through 507.
Attempts were also made to de-protect the pro-ligand in a dichloromethane and pivalic acid mixture and then to remove the dichloromethane to see whether the de-protected ligand would remain dissolved in pivalic acid, but this approach was also unsuccessful. Attempts to form linked wheel systems in an identical fashion using the ruthenium and rhenium complexes 509 and 511 were also unsuccessful. There is a possibility that the use of alternative carboxylic acids as solvents could enable the reaction by dissolving the protected starting material.

5.3.1.3 Coordination chemistry of wheels through their templates

The wheel pro-ligand considered in this section of the chapter 514 was synthesised by a literature preparation.\textsuperscript{62} Previous studies using such wheels as bulky pyridyl ligands have only involved coordination chemistry to first row transition metals under very mild conditions. Therefore, we chose to investigate the stability of these wheels under more severe conditions, such as refluxing in high boiling solvents, in order to be able to form complexes of 2nd and 3rd row transition metals.

A 2.5-fold excess of 514 was reacted with Re\textsuperscript{I}Cl(CO)\textsubscript{5} in refluxing toluene for 24 h (Scheme 31). The product was isolated by column chromatography, eluting with toluene, and a small amount of diffraction-quality crystals were grown by gentle evaporation of a pentane/acetone solution. The crystal structure obtained shows that the linked wheel system 516 is produced and that the wheels are mutually trans, giving a rarely seen meridional \{Re\textsuperscript{I}(CO)\textsubscript{3}\}\textsuperscript{+} core. This is the first reported example of a complex mer-Re\textsuperscript{I}Cl(CO)\textsubscript{3}(L)\textsubscript{2} where L is attached through a pyridyl group, but there are a few related examples in the literature where L is a bulky phosphine.\textsuperscript{128,129,130}

![Scheme 31. Reaction of 514 with Re\textsuperscript{I}Cl(CO)\textsubscript{5}.]
Characterisation data, obtained on the product before crystallisation, suggests that the primary product may in fact be the mono-substituted derivative 515. The IR spectrum and elemental analysis of the powder suggests that it is in fact a rhenium complex containing only one wheel. However, some of the linked wheel system was synthesised. It is believed that approximately 4% of the product formed crystals, and Table 8 shows that the elemental analysis agrees with this allowance.

Table 8. Elemental Analysis of Product from Reaction of 514 with Re\textsuperscript{I}Cl(CO)\textsubscript{5}.

<table>
<thead>
<tr>
<th></th>
<th>Found</th>
<th>515</th>
<th>516</th>
<th>515 + 4% 516</th>
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<td>2647.54</td>
<td>4965.36</td>
<td>2846.15</td>
</tr>
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<td>Re</td>
<td>6.69</td>
<td>6.99</td>
<td>3.75</td>
<td>6.80</td>
</tr>
</tbody>
</table>

Because Ru polypyridyl complexes are the archetypal and most well-studied redox-active MLCT chromophores, a reaction was attempted between \textit{cis}-Ru(bpy)\textsubscript{2}Cl\textsubscript{2}•2H\textsubscript{2}O and 514 (Scheme 32).

Scheme 32. Attempted synthesis of a ruthenium-based linked wheel system.

Following a well established synthetic approach,\textsuperscript{92} \textit{cis}-Ru(bpy)\textsubscript{2}Cl\textsubscript{2}•2H\textsubscript{2}O was reacted with silver triflate in ethanol for 3 hours, removing the chloride ligands and forming an intermediate complex with weakly coordinating triflate ligands. This step is driven by
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The formation of the highly insoluble AgCl, which was removed from the reaction mixture by filtration before the addition of 514. The evidence available, suggests that when 514 is used the product contains only one wheel. This seems logical as the wheel-bearing ligands are expected to coordinate cis to each other as trans-\{Ru(bpy)\}_2^{2+} complexes are very unstable and undergo isomerism back to their cis forms when exposed to light. Considering the bulk and steric restraints of these wheels and their behaviour in the previously mentioned rhenium example, it seems possible that the system may be too sterically hindered to allow a cis arrangement of the wheels.

![Scheme 33. Attempted synthesis of a rhenium-containing wheel system.](image)

Two reactions have been attempted to synthesise a compound of the general formula \( \text{fac-[Re([PyCH}_2\text{NH}_2\text{Et}][\text{Cr}7\text{NiF}_8(\text{O}_2\text{CCMe}_3)_{16}](\text{CO})_3(bpy))]^{+} \) using 514 (Scheme 33). The first method (A) is an approach commonly found in the literature\(^{131,132}\) and has two steps, involving firstly the formation of \( \text{fac-[Re(MeCN)(CO)}_3(bpy)]^{+}[\text{CF}_3\text{SO}_3]^{133} \) as an intermediate by reacting \( \text{fac-ReCl(CO)}_3(bpy) \) with silver triflate in acetonitrile. The MeCN complex is isolated and characterised before being added to two equivalents of 514 and refluxed in THF for 4 hours. This reaction proved to be unsuccessful and one of the possible problems is the use of THF as a solvent; previous studies have shown that the wheel in 514 begins to disintegrate upon sitting in THF for a long period of time. Upon refluxing it is likely that degradation will occur more quickly.

As a result and further to discussions with Dr Mike Coogan, a second more direct method (B) was applied, using toluene as a solvent in the knowledge that linked wheel systems have been synthesised previously in refluxing toluene solutions. In this method, silver
tetrafluoroborate was used as a catalyst and $\text{514}$ was added at the start. However, this approach also proved unsuccessful. In both cases there is evidence to suggest that upon complexation of the template to the rhenium centre loss of the wheel occurs. This is not a wholly unexpected result as the wheels are only attached to the template through hydrogen bonds, which could break under severe conditions.

5.3.1.4 Synthesis of Cr$_7$Co analogues

One major problem with using heterometallic wheels with a ground state spin is that certain techniques, such as $^1$H NMR spectroscopy become unusable as the slow relaxation of the paramagnetic metal centres shifts and broadens the spectra beyond recognition. This means that in general compounds cannot reasonably be identified without obtaining crystal structures. One possible way to deal with this obstacle is to synthesise a series of Cr$_7$Co analogues of the wheel derivatives that are used as ligands. The fast relaxation time of cobalt(II) counteracts the slow relaxation of chromium(III) and thus proton signals can be identified, even though certain features of the spectrum would be missing, such as proton-proton couplings. This technique has already been used with simple symmetrical Cr$_7$Co analogues$^{35}$ hence it was decided to investigate whether this would be a suitable characterisation tool for non-symmetrical wheels and linked wheel systems.

The Cr$_7$Co analogue of $\text{514}$ was synthesised by heating chromium trifluoride and 4-(ethylaminomethyl)pyridine in pivalic acid at 140 °C for 5 hours before adding \{CoPiv$_2$\}_n and increasing the temperature to 160 °C for a further 24 hours (Scheme 34). The impure product [C$_8$H$_5$CH$_2$NH$_2$Et][Cr$_7$CoF$_8$(O$_2$CCMe$_3$)$_{16}$], $\text{517}$ was reacted with [Cu(Piv)$_4$(HPiv)$_2$] to give pure Cu$_2$(O$_2$CCMe$_3$)$_4$([C$_8$H$_5$CH$_2$NH$_2$Et][Cr$_7$CoF$_8$(O$_2$CCMe$_3$)$_{16}$])$_2$, $\text{518}$. This process removed any “horseshoe” clusters that may have been present. The Cu dimer assembly was then reacted with acetylacetone to release the wheels from the complex and give $\text{517}$ in its pure form. Both $\text{517}$ and $\text{518}$ have been characterised using elemental analysis, paramagnetic NMR spectroscopy, mass spectrometry and X-ray crystallography. Crystals were grown by evaporation of a pentane/acetone solution.
Scheme 34. Synthetic route to 517.

5.3.1.5 Synthesis of a new “substitution molecule”

Scheme 35. Synthesis of a novel substitution molecule.

Based on preliminary studies with the two types of derivatised wheel, with the coordinating group in the template or substituted directly into the backbone of the wheel, it appears that the latter are the most useful. Besides iso-nicotinic acid, a series of other compounds have been identified as potential candidates for substitution into the wheel.
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Compounds with extended structures may allow more space between a coordinated metal centre and the wheel, facilitating the preparation of more highly-substituted derivatives.

One compound synthesised for this purpose, but ultimately not further used is \( N \)-\((3\text{-carboxyethyl})\)-4,4'\'-bipyridine hexafluorophosphate, \( 519 \). This was synthesised using a method identical to that used for \( 502 \), but using 3-bromopropionic acid in place of 2-bromoethylamine hydrobromide (Scheme 35). The only other alteration was that purification was effected by metathesis to the \( \text{PF}_6^- \) salt by dissolving the bromide salt in a minimum amount of water and then adding aqueous ammonium hexafluorophosphate. This product has been characterised using \(^1\text{H NMR spectroscopy} and elemental analysis. However, this compound was not further used because other avenues of investigation proved more attractive. \( 519 \) is a salt and it was decided to focus on charge-neutral species for substitution into the wheels as to use a charged carboxylic acid may have large and undesirable effects on the solubility of the resulting wheel-based pro-ligand.

5.3.2 Analysis of \(^1\text{H NMR Spectroscopy}\)

The bromide salts of the three organic quaternised 4,4'\'-bipyridine derivatives were characterised by \(^1\text{H NMR spectroscopy} in \text{D}_2\text{O}. The spectrum of \( 501 \) can be seen in Figure 87.

This compound is symmetric, so the spectrum shows only four signals relating to different environments, two associated with the ethylamine chain and two associated with the bipyridine. The two doublet signals associated with the bipyridine ring are shifted downfield due to quaternisation and appear between 8.5 and 9.2 ppm. No peak is observed for the terminal amine groups, which is to be expected as the protons attached to the nitrogen can undergo rapid exchange with the \( \text{D}_2\text{O} \) solvent.
The \(^1\)H NMR spectrum of the monoquaternised product, 502, can be seen in Figure 88. The two triplet signals associated with the ethylamine chain are in positions virtually identical to those for 501, but there are now four peaks in the aromatic region because the compound is no longer symmetrical. The most deshielded peak at 8.96 ppm can be assigned to the two protons on the carbon adjacent to the quaternised nitrogen. The spectrum for the monoquaternised bpe derivative 503 is very similar with the presence of two additional doublet signals at 7.66 and 7.47 ppm associated with the ethylene unit, the coupling constant of 16.4 Hz confirming the E-geometry.

The \(^1\)H NMR spectrum of 519 is as expected and very similar to that of the amine-based compound 502, with two doublets assigned to the ethylene chain and four doublets assigned to the bpy fragment. As in 502 there is no peak associated with the terminal proton of the chain due to rapid exchange with the deuterium of the solvent.
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Figure 88. $^1$H NMR Spectrum of 502 in D$_2$O.

The $^1$H NMR spectrum of the pro-ligand 506 (Figure 89) was run in deuterated DMF due to solubility issues. A large singlet at 1.58 ppm arises from the 18 protons of the $^3$Bu groups, and two triplets between 3.53 and 3.73 ppm are due to the two CH$_2$ groups. Two broad peaks at 7.15 and 9.25 ppm are attributable to the amide protons, that adjacent to the pyridyl rings being shifted well into the aromatic region. Three other peaks are observed in the aromatic region due to the pyridyl rings.

Figure 89. The $^1$H NMR Spectrum of the pro-ligand 506 in $d_6$-DMF.
The spectrum of the Re complex 511 is very similar to that of 506, but a direct comparison of the spectra in deuterated DMF shows that most of the signals for 511 are further deshielded when compared with those of 506, especially the bpy signals which are deshielded by as much as 0.41 ppm. This observation can be attributed to the electron-withdrawing influence of the fac-ReCl(CO)₃ unit. The signal for the remote tBu groups is predictably insensitive to complexation.

The ¹H proton NMR spectrum of 511 has also been run in deuterated acetone in order to allow comparisons with the spectrum of 508 (Figure 90). For the latter it is harder to assign particular peaks to the BOC protected ligand as the aromatic region is much more complicated. In general, however, it can be observed that once again the peaks associated with the Re complex 511 are more deshielded.

Figure 90. The ¹H NMR spectrum of 511 in acetone. Inset: The bpy region of the spectrum of 508.

5.3.3 Paramagnetic NMR studies

Paramagnetic ¹H NMR studies are a potential diagnostic tool to identify wheel systems in the solution state when there is no crystal structure available. It may be possible to
identify how many wheels are attached to a metal centre by using the relative integrals in the spectrum. The two novel Cr₇Co systems described in this chapter have hence been studied using this technique. The full spectrum of 517 can be seen in Figure 91. The spectrum of 518 was too broad to yield any reasonable data due to the extra paramagnetism of the copper centres in the template, so it will not be discussed further.

The peaks are, as expected, broader than those normally seen in a ¹H NMR experiment and are spread over a far wider range than the typical 0–10 ppm, but the pivalate protons are five bonds away from the paramagnetic centres and therefore have signals sharp enough to be identified. Peaks assigned to the pivalates occur between 15 and −3 ppm and those assigned to the template lie between −5 and −35 ppm, showing that the template remains in the cavity in solution.

![Figure 91. Paramagnetic ¹H NMR Spectrum of 517 in CDCl₃. Insert: expanded pivalate region.](image)

As expected for wheels with asymmetric templates, this spectrum shows twelve peaks corresponding to the pivalate groups. Four of these peaks are assigned to the equatorial pivalates, each corresponding with two groups. This observation can be accounted for by
the fact that the equatorial pivalates do not feel any effect from the template and are only differentiated by their proximity to the hetero-metal. The other eight peaks can be assigned to the axial pivalates, each one different as they are affected by their location with respect to the hetero-metal and to the template. A small but sharp signal can be seen at 7.26 (*) corresponding to the solvent used, CDCl$_3$. The template region is very busy and hard to assign due to the number and size of the peaks.

5.3.4 IR Studies

While NMR spectroscopy shows no difference in structure between 505 and 506, IR spectroscopy does allow these two compounds to be distinguished. A clear difference is seen in the N–H regions, with 506 showing two distinct bands at 3348 and 3298 cm$^{-1}$ corresponding with the presence of two different secondary amide groups. The spectrum of 505 shows a shift to a lower wavenumber, as expected for an absorption caused by an amine instead of an amide N–H bond, and also a much broader absorption band in that region, which can be attributed to the increasing H-bonding ability of the primary protonated amine. Other differences are observed in the C=O (amide) stretching region, 506 showing two bands between 1690 and 1630 cm$^{-1}$, whereas 505 shows only one band.

Figure 92. IR spectra of 505 and 506.
in this region and one band at 1560 cm\(^{-1}\) correlating to the carboxylate salt region. These differences in the spectra can be seen in Figure 92.

As can be seen in Figure 93, the IR spectrum of 511 shows three strong carbonyl stretching absorptions at 2026, 1919 and 1880 cm\(^{-1}\) which are characteristic of a \(\text{fac-}\{\text{Re}^{I}(\text{CO})_{3}\}\)\(^{+}\) unit. A broad peak corresponding to the two amide carbonyl groups in the ligand is also present at 1656 cm\(^{-1}\), in contrast with the two sharp \(\nu(\text{C}=\text{O})\) peaks in the spectrum of the pro-ligand.

![Figure 93. IR spectrum of 511.](image)

5.3.5 UV-VIS studies

The UV-VIS spectra of 509 and 511 are shown in Figure 94 and the corresponding data are collected in Table 9. As expected, broad bands due to MLCT transitions can be seen in the region of approximately 400–500 nm, and other bands are observed to higher energies, attributable primarily to intraligand \(\pi-\pi^{*}\) transitions. These values do not differ greatly from the literature values obtained for the mono-armed Ru system.\(^{118}\)
Figure 94. UV-VIS spectra of 509 and 511 in acetonitrile at 293 K.

Table 9. UV-VIS data for 509 and 511 in acetonitrile.

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<tbody>
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</tr>
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</table>

5.3.6 Electrochemistry

The ruthenium complex 509 was studied using cyclic voltammetry in 0.1 M solution of [NBu<sub>4</sub>][PF<sub>6</sub>] in acetonitrile. A representative voltammogram can be seen in Figure 95. A fully reversible process is observed at \( E_{1/2} = 1.41 \) V vs. Ag–AgCl (\( ΔE_p = 60 \) mV), as expected for such complexes, relating to the Ru<sup>III/II</sup> oxidation. In addition, three quasireversible processes are observed for the bpy ligands at \( E_{1/2} = -1.05 \) V, -1.42 V and -1.65 V (\( ΔE_p = 70 \) mV, 55 mV and 55 mV, respectively).
5.3.7 Crystal structure determination

5.3.7.1 Organic Molecules

Crystals of 505 were grown by vapour diffusion of diethyl ether into DMSO and a representation of the molecular structure can be seen in Figure 96. Only one cation is present in this representation, although there are two cations in the salt. The position of H13 has been refined so it is possible to calculate the distance between it and O2. This distance is relatively short and the angle between N2–H13….O2 is 166°, indicating that there is hydrogen bonding between the molecules. This is as is expected and helps to explain the crystal packing arrangement (Figure 97), as well as the broadened N–H peaks seen in the IR spectrum.
Figure 96. Representation of the molecular structure of 505.

Figure 97. Crystal packing diagram of 505.

Crystals of 502.(PF$_6$)$_2$ and 502.(BPh$_4$)$_2$ were grown by vapour diffusion of diethyl ether into acetone solutions at room temperature. Representations of the molecular structures can be seen in Figures 98 and 99 respectively. Interestingly, in both cases the product has reacted with the acetone solvent to form a Schiff base derivative. This observation is unexpected because it is believed that both terminal amine groups were protonated before crystallisation and therefore would not have had a free lone pair to donate in the Schiff base formation. A key difference between the two crystal structures is that the nitrogen atom of the imine group is still protonated in the crystal of the PF$_6^-$ salt but no longer protonated in the crystal of the BPh$_4^-$ salt.
The counter-anion affects the crystallisation properties greatly, with the PF₆⁻ salt adopting the orthorhombic $Pca2_1$ space group and the BPh₄⁻ salt adopting the monoclinic
Chapter 5 – Linked Wheels through Hydrogen Bonds

\[ P2_1/n \] space group. The counter-anion may also have determined whether the amine remained protonated following the condensation reaction with acetone. Other selected crystallisation data for the three salts can be seen in the Appendix.

### 5.3.7.2 Crystal structures of wheels and linked wheel systems

Crystals of 516 were obtained by evaporation of a 3:2 \( n \)-pentane/acetone solution at room temperature. The X-ray diffraction data were obtained at the University of Manchester by Dr Elizabeth Harper and Dr John Fielden. This crystal structure has not been finalised due to the severe disorder of the templating molecule, but the partially completed molecular structure can be seen in Figure 100.

![Unfinished Representation of the Molecular Structure of 516.](image)

This crystal structure clearly shows that the wheels are located trans to each other, meaning that the three carbonyl groups are coordinated in a meridional fashion around the rhenium centre. The rhenium itself, shown in pink, is disordered over two sites, meaning that the attached pyridyl units are also disordered. The chloride ligand is disordered over four sites, with each site having \( \frac{1}{4} \) occupancy, and the carbonyl ligands
are disordered over four sites each having $\frac{1}{4}$ occupancy, giving a ratio of one chloride to three carbonyl ligands.

The crystal structures of 517 and 518, both derived from data obtained at Daresbury, have not been fully solved as previous Cr7Ni analogous structures have already been finalised. From the raw data, it can be stated that the desired products have been synthesised. Both crystals were grown by evaporation of $n$-pentane/acetone solutions at room temperature.
5.4 CONCLUSIONS

A selection of purely organic molecules and metal complexes with potential applications as switchable linkers has been synthesised and characterised. However, attempts to use these as templates for the wheels, either in a direct synthetic method or in a template substitution reaction, have thus far proved unsuccessful. The direct synthesis method was unsuccessful mainly due to solubility factors; none of the templating molecules synthesised are soluble in pivalic acid, which was used as the reaction solvent. The template substitution reactions were unsuccessful, probably as a result of the wheels becoming detached from their templates.

While it has been proven that linking wheels directly through their backbones is much more effective, two novel indirectly-linked wheel systems have been synthesised, including a rare example of a mer-ReCl(CO)$_3$(L)$_2$ complex.

Ultimately it was concluded that linking wheels together through hydrogen bonds was not a particularly fruitful avenue of investigation and thus more effort was focused on linking the wheels together directly.
Conclusions
Conclusions

The primary aim of this project was to further develop the series of linked wheel systems. This has been achieved, on the whole, with great success but much further work is required on the study of these linked wheel systems to determine how interactions can be affected between the wheels and if it is possible to observe any magnetic switchable effects occurring with some of the MLCT/redox active links that were synthesised towards the end of the project.

A number of accomplishments have been achieved synthetically. It has been proven that selective substitution of functionalised carboxylates into the wheels is possible using a variety of carboxylates, 203 - 210. These can then be used as ligands in complexation to metal centres. The series of first row transition metal complexes has been expanded using these novel wheel based ligands using facile crystallisation techniques (211- 219). These compounds are not expected to show switchable properties, but are anticipated to help understanding of how best to enable interaction between wheels. Certain complexes have been measured by EPR, however many more need to be studied in order to obtain a fuller picture on how best to interact the wheels.

The original wheel based ligand, containing an iso-nicotinate group has also been used as a bulky ligand in 2\textsuperscript{nd} and 3\textsuperscript{rd} row transition metal complexes, undergoing more harsh reaction conditions and column chromatography (220 – 222). Despite no crystallographic evidence, it is strongly believed that metal complex links between purple wheels have been achieved using the qpy, bbpe and pytpy ligands (312 – 315). While initial UV-VIS and cyclic voltammetry measurements have been used as characterisation techniques, there are suggestions that some of these complexes may demonstrate switchable properties. Further studies must be done to verify this, including time-resolved EPR spectroscopy to witness if the excited state is long lived enough to affect the interaction occurring between the wheels.

The series of organic linked wheel systems has been expanded, both by using organic linkers to join together two (or three) purple wheels (301 – 304). Organic links have also been utilised to form a novel series of mixed green-purple linked wheel systems, with variations being different carboxylate ligands acting as linkers and different divalent
metals being used in the purple wheel backbone (401 – 414). EPR studies have been undertaken on these systems showing some interesting results. There is evidence that the carboxylate affects the strength of interaction between the wheels, with the strongest interaction occurring using the iso-nicotinate ligand and the weakest using the 4-pyridazine carboxylate ligand, while use of a Cr7Zn purple wheel effectively switches off interaction between the wheels.

Synthetically it has been proven that attempting to link wheels together indirectly is the most difficult method. The primary problem that arose during the attempts at direct synthesis of linked wheel systems was the lack of solubility of the templating molecules in the carboxylic acid solvent. When attempting to form linked wheel systems using a pre-made wheel containing a pendant pyridyl group, two difficulties became clear. The first was the bulk of the ligand, which was too great to enable any coordination cis to another wheel as displayed upon trans coordination to a ReI metal centre, leaving the carbonyl ligands in a kinetically unfavourable mer arrangement (516). This meant the synthesis of any complex where the wheels needed to be in a cis arrangement, such as a RuII(bpy)2(L)2 type complex, became impossible. The second problem, was the ability of the wheel to “slip off” the template during more harsh reaction conditions, leaving the desired product minus the wheel. Given the lack of entanglement of wheels with these systems this is the least important avenue to be further investigated.

The areas of particular interest for future work are the potentially switchable systems, although it must first be established whether either the MLCT or redox activity in the previously synthesised systems do prove to alter any interaction between the wheels. Further suitable synthetic targets have been outlined at the end of each of the respective chapters, considering only RuII and ReI complexes, however the possibilities of transition metal switchable systems are enormous, and should be considered should this project be moved forward.
Appendix
Table 10. Crystallographic data and refinement details of solved crystal structures.

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<tr>
<th>Formula</th>
<th>211 ( \text{C}<em>{193} \text{H}</em>{344} \text{Cr}<em>{14} \text{Cu}</em>{2} \text{F}_{16} )</th>
<th>213 ( \text{C}<em>{219} \text{H}</em>{376} \text{Cr}<em>{14} \text{Cu}</em>{2} \text{F}_{16} )</th>
<th>215 ( \text{C}<em>{196} \text{H}</em>{350} \text{Cr}<em>{14} \text{Cu}</em>{1} \text{F}<em>{16} \text{N}</em>{6} )</th>
<th>217 ( \text{C}<em>{182} \text{H}</em>{310} \text{Cr}<em>{14} \text{F}</em>{28} \text{N}_{6} )</th>
<th>219 ( \text{C}<em>{311} \text{H}</em>{564} \text{Cr}<em>{13} \text{Cu}</em>{21} \text{F}<em>{24} \text{Fe}</em>{3} \text{N}<em>{12} \text{Ni}</em>{2} \text{O}_{112} )</th>
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<td>Orthorhombic</td>
<td>Triclinic</td>
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<td>( P \bar{1} )</td>
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