Synthesis and Reactivity of Group 12 β-Diketiminate Coordination Complexes

By

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Abstract

The variable β-diketiminate ligand poses as a suitable chemical environment to explore unknown reactivity and functionality of metal centres. Variants on the β-diketiminate ligand can provide appropriate steric and electronic stabilization to synthesize a range of β-diketiminate group 12 metal complexes. This project aimed to explore various β-diketiminate ligands as appropriate ancillary ligands to derivatise group 12 element complexes and investigate their reactivity.

A β-diketiminato-mercury(II) chloride, \([o-C_6H_4{C(CH_3)=N-2,6-iPr_2C_6H_3}{NH(2,6-iPr_2C_6H_3)}]HgCl\), was synthesized by addition of \([o-C_6H_4{C(CH_3)=N-2,6-iPr_2C_6H_3}{NH(2,6-iPr_2C_6H_3)}]Li\) to mercury dichloride. Attempts to derivatise the β-diketiminato-mercury(II) chloride using salt metathesis reactions were unsuccessful with only β-diketiminate ligand degradation products being observed in the \(^1\)H NMR.

A β-diketiminato-cadmium chloride, \([CH{(CH_3)CN-2,6-iPr_2C_6H_3}2]CdCl\), was derivatized to a β-diketiminato-cadmium phosphanide, \([CH{(CH_3)CN-2,6-iPr_2C_6H_3}2]Cd P(C_6H_{11})_2\), via a lithium dicyclohexyl phosphanide and a novel β-diketiminato-cadmium hydride, \([CH{(CH_3)CN-2,6-iPr_2C_6H_3}2]CdH\), via Super Hydride. Initial reactivity studies of the novel cadmium hydride with various carbodiimides yielded a β-diketiminato-homonuclear cadmium-cadmium dimer, \([CH{(CN-2,6-iPr_2C_6H_3}2]Cd\), which formed via catalytic reduction of the cadmium hydride. Attempts to synthesize an amidinate insertion product via a salt metathesis reaction or a ligand exchange reaction proved unsuccessful but a novel cadmium amidinate, \([{CH(N-C_6H_{11})_2}{CH(N-C_6H_{11})(N(H)-C_6H_{11})}]Cd\), was synthesized from addition of dicyclohexyl formamidine to bis-hexamethyldisilazane cadmium.

A β-diketiminato-zinc(II) bromide, \([o-C_6H_4{C(CH_3)=N-2,6-iPr_2C_6H_3}{NH(2,6-iPr_2C_6H_3)}]ZnBr\), was synthesized by addition of \([o-C_6H_4{C(CH_3)=N-2,6-iPr_2C_6H_3}{NH(2,6-iPr_2C_6H_3)}]Li\) to zinc dibromide. The β-diketiminato-zinc(II) bromide was derivatized to a variety of complexes (including amides and phosphanides) by a salt metathesis reaction. Chalcogen addition reactions were performed from \([o-C_6H_4{C(CH_3)=N-2,6-iPr_2C_6H_3}{NH(2,6-iPr_2C_6H_3)}]ZnP(C_6H_{11})_2\] to produce double addition products from sulfur, selenium and
tellurium. Chalcogen addition reactions from \([o-C_6H_4(C(CH_3)=N-2,6-^{1}Pr_2C_6H_3)NH(2,6-^{1}Pr_2C_6H_3)]ZnP(C_6H_5)_2\) produced a double addition product for selenium and a β-diketiminato-zinc(II) tellunoite bridged dimer, \([o-C_6H_4(C(CH_3)=N-2,6-^{1}Pr_2C_6H_3)NH(2,6-^{1}Pr_2C_6H_3)]ZnTe\), from tellurium. A total of 14 compounds were characterized via X-ray diffraction. Photoluminescence studies of the β-diketiminato-zinc(II) compounds were conducted where it was proposed that an electron transfer from the lone pair on the hetero-atom influenced the quantum yield and fluorescence intensities.
Acknowledgements

Firstly, I would like to express my gratitude to my supervisor Dr. Robin Fulton for her continuous support of my research, her patience and consistent motivation. She constantly allowed this thesis to be my own work, but always steered me in the right the direction.

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<table>
<thead>
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<th>Symbol</th>
<th>Description</th>
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<tbody>
<tr>
<td>δ</td>
<td>Chemical shift (ppm)</td>
</tr>
<tr>
<td>λ</td>
<td>Wavelength</td>
</tr>
<tr>
<td>λ&lt;sub&gt;Max&lt;/sub&gt;</td>
<td>Wavelength of maximum absorption</td>
</tr>
<tr>
<td>NMR</td>
<td>Nuclear magnetic resonance</td>
</tr>
<tr>
<td>1H NMR</td>
<td>Hydrogen nuclear magnetic resonance</td>
</tr>
<tr>
<td>13C NMR</td>
<td>Carbon nuclear magnetic resonance</td>
</tr>
<tr>
<td>31P NMR</td>
<td>Phosphorus nuclear magnetic resonance</td>
</tr>
<tr>
<td>Å</td>
<td>Angstrom</td>
</tr>
<tr>
<td>Ar (images)</td>
<td>Commonly used aryl group 2,6-diisopropylphenyl (Dipp)</td>
</tr>
<tr>
<td>Ar (characterization)</td>
<td>Aryl or aromatic region</td>
</tr>
<tr>
<td>Am</td>
<td>Amidinate</td>
</tr>
<tr>
<td>Approx.</td>
<td>Approximately</td>
</tr>
<tr>
<td>BDI</td>
<td>β-diketiminate</td>
</tr>
<tr>
<td>BDI&lt;sub&gt;Dipp&lt;/sub&gt;</td>
<td>β-diketiminate ligand, [CH{((CH&lt;sub&gt;3&lt;/sub&gt;)CN-2,6-&lt;sup&gt;i&lt;/sup&gt;Pr&lt;sub&gt;2&lt;/sub&gt;C&lt;sub&gt;6&lt;/sub&gt;H&lt;sub&gt;3&lt;/sub&gt;}]&lt;sup&gt;-&lt;/sup&gt;</td>
</tr>
<tr>
<td>Ar-BDI&lt;sub&gt;Dipp&lt;/sub&gt;</td>
<td>Aryl incorporated β-diketiminate ligand, [o-C&lt;sub&gt;6&lt;/sub&gt;H&lt;sub&gt;4&lt;/sub&gt;{C(CH&lt;sub&gt;3&lt;/sub&gt;)=N-2,6-&lt;sup&gt;i&lt;/sup&gt;Pr&lt;sub&gt;2&lt;/sub&gt;C&lt;sub&gt;6&lt;/sub&gt;H&lt;sub&gt;3&lt;/sub&gt;}{NH(2,6-&lt;sup&gt;i&lt;/sup&gt;Pr&lt;sub&gt;2&lt;/sub&gt;C&lt;sub&gt;6&lt;/sub&gt;H&lt;sub&gt;3&lt;/sub&gt;)}]&lt;sup&gt;-&lt;/sup&gt;</td>
</tr>
<tr>
<td>br</td>
<td>Broad signal</td>
</tr>
<tr>
<td>C&lt;sub&gt;6&lt;/sub&gt;D&lt;sub&gt;6&lt;/sub&gt;</td>
<td>Deuterated benzene</td>
</tr>
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<td>Cy</td>
<td>Cyclohexyl</td>
</tr>
<tr>
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<td>Doublet</td>
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<td>Doublet of doublets</td>
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<td>Doublet of doublet of doublets</td>
</tr>
<tr>
<td>DCM</td>
<td>Dichloromethane</td>
</tr>
<tr>
<td>Dipp</td>
<td>2,6-diisopropylphenyl</td>
</tr>
<tr>
<td>E</td>
<td>Chalcogen (sulfur, selenium or tellurium)</td>
</tr>
<tr>
<td>eq</td>
<td>Equation</td>
</tr>
<tr>
<td>Eq.</td>
<td>Equivalence</td>
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<tr>
<td>Et</td>
<td>Ethyl</td>
</tr>
<tr>
<td>Et&lt;sub&gt;2&lt;/sub&gt;O</td>
<td>Diethyl ether</td>
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<tr>
<td>Formamidine</td>
<td>Fo</td>
</tr>
<tr>
<td>&lt;sup&gt;i&lt;/sup&gt;Pr</td>
<td>iso-propyl substituent</td>
</tr>
<tr>
<td>i</td>
<td>Ipso- position</td>
</tr>
</tbody>
</table>
**IR**  Infrared spectroscopy

**HMDS**  Hexamethyldisilazane

**Hex**  Hexane

**J**  Scalar coupling constant (Hz)

**LDA**  Lithium diisopropyl amide

**L<sub>n</sub>**  An unspecified number of an unspecified ligand

**m**  Multiplet

**m**  Meta- position

**Me**  Methyl

**mes**  Mesitylene group

**"Bu**  n-butyl substituent

**o**  Ortho- position

**p**  Para- position

**Ph**  Phenyl

**ppm**  Parts per million

**pz**  Pyrazole ring

**R**  An unspecified substituent

**s**  Singlet

**t**  Triplet

**td**  Triplet of doublets

**terp**  Terphenyl ligand

**tol**  Toluene

**THF**  Tetrahydrofuran

**TMS**  Trimethylsilyl

**Tp**  Trispyrazol borate ligand

**UV/vis**  Ultraviolet-visible spectroscopy

**v**  Wavenumbers (cm<sup>-1</sup>)
Nomenclature

In this thesis, most of the compounds are supported by a bulky β-diketiminate ligand. Various β-diketiminato-group 12 metal complexes containing various substituents were synthesized. There are several ways of naming these compounds; the nomenclature generally used in this thesis is set out as below. Often the metal is omitted from the abbreviated name when clarification is not necessary.

Table i: Nomenclature of the compounds synthesized in this thesis, where \( L = \text{BDI}_{\text{Dipp}} \) or \( \text{Ar-BDI}_{\text{Dipp}} \), \( \text{Metal} = M = \text{Zn, Cd or Hg} \).

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<thead>
<tr>
<th>Structure</th>
<th>Nomenclature</th>
<th>Abbreviated name</th>
</tr>
</thead>
<tbody>
<tr>
<td>[LMCl]</td>
<td>( \beta )-diketiminato-metal(II) chloride</td>
<td>Metal chloride</td>
</tr>
<tr>
<td>[LMBr]</td>
<td>( \beta )-diketiminato-metal(II) bromide</td>
<td>Metal bromide</td>
</tr>
<tr>
<td>[LMI]</td>
<td>( \beta )-diketiminato-metal(II) iodide</td>
<td>Metal Iodide</td>
</tr>
<tr>
<td>[LMBr(_2)Li(S)(_n)]</td>
<td>( \beta )-diketiminato-metal(II) dibromide lithium salt</td>
<td>Metal dibromide lithium salt</td>
</tr>
<tr>
<td>[LMHMDS]</td>
<td>( \beta )-diketiminato-metal(II)</td>
<td>Metal HMDS</td>
</tr>
<tr>
<td>[LMN(H)Dipp]</td>
<td>( \beta )-diketiminato-metal(II) 2,6-diisopropylphenyl amide</td>
<td>Metal anilido</td>
</tr>
<tr>
<td>[LMH]</td>
<td>( \beta )-diketiminato-metal(II) hydride</td>
<td>Metal hydride</td>
</tr>
<tr>
<td>[LM](_2)</td>
<td>( \beta )-diketiminato-homonuclear metal-metal dimer</td>
<td>Homonuclear M-M dimer</td>
</tr>
<tr>
<td>[LMPCy(_2)]</td>
<td>( \beta )-diketiminato-metal(II) dicyclohexylphosphanide</td>
<td>Metal dicyclohexylphosphanide</td>
</tr>
<tr>
<td>[LMPPh(_2)]</td>
<td>( \beta )-diketiminato-metal(II) diphenylphosphanide</td>
<td>Metal diphenylphosphanide</td>
</tr>
<tr>
<td>[LM(S)(_2)PCy(_2)]</td>
<td>( \beta )-diketiminato-metal(II) dicyclohexylphosphinodithioate</td>
<td>Metal dicyclohexylphosphinodithioate</td>
</tr>
<tr>
<td>[LM(Se)(_2)PCy(_2)]</td>
<td>( \beta )-diketiminato-metal(II) dicyclohexylphosphinodiselenoate</td>
<td>Metal dicyclohexylphosphinodiselenoate</td>
</tr>
<tr>
<td>[LM(Se)PCy(_2)]</td>
<td>( \beta )-diketiminato-metal(II) dicyclohexylphosphinoselenoite</td>
<td>Metal dicyclohexylphosphinoselenoite</td>
</tr>
<tr>
<td>Structure</td>
<td>Nomenclature</td>
<td>Abbreviated name</td>
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<tr>
<td>-----------</td>
<td>--------------</td>
<td>-----------------</td>
</tr>
<tr>
<td>[LM(Te)₂PCy₂]</td>
<td>β-diketiminato-metal(II)</td>
<td>Metal dicyclohexylphosphinoditellunoate</td>
</tr>
<tr>
<td>[LM(Se)₂PPh₂]</td>
<td>β-diketiminato-metal(II)</td>
<td>Metal diphenylphosphinodiselenoate</td>
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<tr>
<td>[LM(Se)PPh₂]</td>
<td>β-diketiminato-metal(II)</td>
<td>Metal diphenylphosphinosalenoite</td>
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<td>[LM(Te)PPh₂]</td>
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<td>Metal diphenylphosphinoditellunoate</td>
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<td>[LM(Te)PPh₂]</td>
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<td>Metal diphenylphosphinoditellunoate</td>
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<tr>
<td>[LM(Te)₂PPh₂]</td>
<td>β-diketiminato-metal(II)</td>
<td>Metal diphenylphosphinoditellunoate</td>
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<tr>
<td>[LM][₂]Te</td>
<td>β-diketiminato-metal(II)</td>
<td>Metal tellunoite bridged dimer tellunoite bridged dimer</td>
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<tr>
<td>Cy-N=N-Cy</td>
<td>dicyclohexyl formamidine</td>
<td>DC-Fo</td>
</tr>
<tr>
<td>iPr-N=N-iPr</td>
<td>diisopropyl formamidine</td>
<td>DI-Fo</td>
</tr>
<tr>
<td>Cy-N=M-N-Cy</td>
<td>dicyclohexyl amidinate metal(II)</td>
<td>DC-Am metal (M = Li or Cd)</td>
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<tr>
<td>iPr-N=M-iPr</td>
<td>diisopropyl amidinate metal(II)</td>
<td>DI-Am metal (M = Li or Cd)</td>
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<tr>
<td>Cy-N=C=N-Cy</td>
<td>dicyclohexylcarbodiimide</td>
<td>DCC</td>
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<tr>
<td>iPr-N=C=N-iPr</td>
<td>diisopropylcarbodiimide</td>
<td>DIC</td>
</tr>
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</table>
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# Structural List of Newly Synthesized Compounds

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<thead>
<tr>
<th>Abbreviated name</th>
<th>Complex</th>
<th>Structure</th>
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<td>XVI</td>
<td><img src="image" alt="CdCl2" /></td>
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<tr>
<td>Zinc dicyclohexylphosphanide</td>
<td>XXX</td>
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<tr>
<td>Zinc diphenylphosphanide</td>
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<td><img src="image" alt="ZnPPh2" /></td>
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<td>Zinc dibromide lithium salt</td>
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Chapter 1

Background and Literature

1.1 Introduction

1.1.1 Group 12 Elements

Group 12 elements include the three naturally occurring elements zinc (Zn), cadmium (Cd) and mercury (Hg) and the recently synthesized element copernicium (Cn). Despite sharing qualities based on their appearance in the periodic table, each of the three natural group 12 elements differ to each other in chemical reactivity and coordination chemistry.

The natural abundance of zinc is 65 ppm, the low abundancy of cadmium and mercury, 0.1 ppm 0.08 ppm respectively, limits their use in higher biological organisms. Zinc is the only group 12 element that is an essential trace element and after iron is the second most abundant transition metal in the human body. Zinc has been shown to be present in approximately 10% of human proteins and is heavily regulated within the body. This makes zinc the only biologically relevant group 12 element.

The low melting and boiling points of the group 12 elements indicates a weak metallic bond with little valence band and conduction band overlap. This pushes their metallic properties close to the boundary of a metalloid element like the main-group elements gallium (Ga) and germanium (Ge). The predominant oxidation state for group 12 elements in complexes is +2, giving them a d\textsuperscript{10} electronic configuration with other oxidation states being rare. Coordination to substituents only occurs through the s-orbitals, the d-orbitals remain core-like. In these ways the group 12 elements share similarities with main-group elements.

IUPAC (The International Union of Pure and Applied Chemistry) defines a transition metal element “... as an element whose atom has an incomplete d sub-shell” of which mercury is the only culprit with an oxidation state higher than +2 in the rare non-equilibrium conditions
creating mercury (IV) fluoride. Group 12 elements zinc and cadmium share qualities to group 2 elements beryllium (Be) and magnesium (Mg) in atomic radii, ionic radii, electronegativities, binary compound structure and their ability to form complex ions with natural nitrogen and oxygen ligands. The similarity with group 2 elements goes a far as the old IUPAC system naming them as 2-B. The group 12 elements have been given the name outer transition metals in earlier scientific texts due to their differences to transition metals but similarities to group 2 elements.

The ambiguity of the classification and inconsistent qualities for all three naturally occurring group 12 elements calls for an in depth investigation into their synthetic and coordination chemistry. Zinc has been heavily investigated due to its non-toxic qualities and biological relevance. In contrast cadmium and mercury have received less attention. There is therefore a gap in the understanding of zinc, cadmium and mercury’s coordination chemistry.

1.1.2 The β-Diketiminate Ligand

The β-diketiminate ligand, abbreviated as either NacNac or more commonly BDI, can be utilized as an ancillary ligand, supplying both steric and electronic stabilization to a metal centre. β-Diketiminate ligands have been coordinated to almost every available element with exceptions including some heavier transition and main-group metals or rare elements. β-Diketiminate ligands have been used to understand the behaviour of metals and metal coordination in biological environments. The β-diketiminate system represents a perfect chemical environment for an investigation into the coordination of group 12 elements to compare their complexes.

![](image)

**Figure 1:** BDI Substitution sites. Left, bare ligand with delocalized electron. Right, metallocycle formed with metal centre with α, β and γ positions.

The substituents on the backbone of β-diketiminate ligands can be varied to allow fine tuning of the ligand’s electronic and steric properties (Figure 1). At the γ-position, the R³
substituent is typically a hydrogen atom and is referred to as the γ-proton. The π-electrons are delocalized across the backbone when the ligand is coordinated to a metal centre. The coordination of β-diketiminate ligands is commonly through the two nitrogen atoms, forming a metallocycle with the metal centre.

The coordination of BDI to a metal centre results in planar geometry of the ligand due to delocalized π-electrons. The BDI ligand is typically a 4-electron donor, formally through amide and imine donor nitrogens of the ligand. Amsterdam Density Functional Theory Calculations were used to explain the metal’s bonding orbitals which were in plane with the σ-bonds that formed with the ligands nitrogen atoms. With sufficient steric crowding of the BDI substituents, the ligand can adopt a highly puckered boat configuration which has varying amount of non-planarity (Figure 2). The highly puckered boat system allows occupied out of phase π-orbitals from the nitrogen atoms to bond to the metal centre making the BDI ligand a 6-electron donor in a $2\sigma + \pi$ process.

![Figure 2: Left, planar conformation of BDI. Right, puckered boat conformation of BDI. Substitution sites have been removed for clarity.](image-url)
1.2 Project Aims

The aim of this project was to investigate two β-diketiminate ligands as appropriate ancillary ligands for each group 12 element for the synthesis of novel complexes. The two β-diketiminate ligands contain sufficient steric bulk to prevent oligomerisation of the resulting group 12 complexes (Figure 3).

![Figure 3: β-Diketiminate ligands used in this project.](image)

To date, only a handful of cadmium and mercury complexes bearing the β-diketiminate ligand have been synthesised, generally from toxic reagents. As the reactivity of the resulting cadmium and mercury complexes have not been completely investigated, this provided an opportunity to explore unseen novel functionality including the synthesis of hydrides and homonuclear metal-metal (M-M) bonded dimers. Many zinc complexes bearing the β-diketiminate ligand have been synthesized but certain functionalities remain absent in the literature.

1.3 Important Chemical Reactions

Throughout this thesis specific reactions are conducted and discussed. The following are important reaction archetypes and chemical processes that involve the metal complexes and the novel functionalities in this project.

Salt Metathesis Reaction

A salt metathesis reaction is defined as the exchange of bonds between two chemical species or reactants where one of the reactants and one of the products is a salt (AgNO₃ and AgCl, Figure 4). The resulting products share similar bonding associations but contain new substituents.¹⁴ Usually one of the products is a precipitate in solution, often the
salt, leaving the other product dissolved in a solvent. The two products can therefore be separated by a filtration. When a coordinating solvent is used, such as diethyl ether (Et₂O) or tetrahydrofuran (THF), the salt metathesis reaction doesn’t always go to completion resulting in a solvated salt coordinated to one of the reactants.

\[
\text{AgNO}_3 + \text{HCl} \rightarrow \text{HNO}_3 + \text{AgCl}
\]

**Figure 4: A typical salt metathesis reaction followed by an example.**

**Insertion Reaction**

An insertion, or addition reaction, involves the interposition of a molecule or an atom into an existing bond of another molecule (Figure 5). In inorganic chemistry, this type of reaction is used to describe a circumstance where the metal centre (M) remains mostly unchanged with no alterations to the coordination number or the oxidation state. The ligand (R), or pre-existing substituent, can change oxidation state or coordination when a substrate (X-Y) is inserted between the M-R bond. The reaction can be reversible as is often the case when an insertion reaction is a step in a catalytic cycle. The reverse reaction is referred to as an elimination or an extrusion reaction.

\[
\text{M-R + X-Y} \leftrightarrow \text{M-X-Y-R} \leftrightarrow \text{M-X-R}
\]

**Figure 5: Common insertion geometries, 1,2 (top) and 1,1 (below).**

**Hydride Ligand and Reactivity**

A hydride is formally the anionic form of hydrogen, referred to as H⁻. A metal hydride complex or hydride may refer to a compound containing one or more of these hydride ligands. Hydride containing complexes represent catalytic intermediates in various homo- and hetero-geneous catalytic cycles. This includes hydrogenation, hydroformylation, hydrosilylation and hydrodesulfurization.
Spectroscopic characterization of metal hydrides involves multinuclear NMR and IR spectroscopies. Structural characterization is possible with X-ray diffraction by finding residual electron densities in the difference density map and using distortions from neighbouring atoms to indirectly determine the hydrides position. Since hydride ligands lack substantial electron density, only neutron diffraction can accurately measure the metal-hydride bond distance and associated angles.

A variety of different types of coordination environments have been observed for metal hydride complexes including classical (A), non-classical (B), bridging (C,D) and interstitial (E,F) hydrides (Figure 6). Classical refers to a hydride ligand acting as an H⁻ ligand, non-classical refers to dihydrogen coordinating to a metal via a σ-bond, bridging refers to a hydride ligand shared between metal centres and an interstitial hydride refers to an internal hydride typically within a metal cluster.¹⁷⁻¹⁸

![Figure 6: From left to right: Classical (A), non-classical (B), asymmetric bridging (C), symmetric bridging (D) and interstitial (E,F). Other substituents are omitted for clarity.](image)

The reactivity of a classical or terminal metal hydride includes migratory insertion, classical insertion and protonolysis, all of which require the metal complex to lose the hydride ligand. Migratory insertion involves the reversible insertion of a hydride across an already coordinated π-complex, such as an alkene (eq 1).

\[
\text{LnM} - \text{H} \rightarrow \text{LnM} - \text{H} \quad (1)
\]

In classical insertion reactions the hydride acts as a nucleophile. An insertion reaction may involve a heteroallene containing a central carbon (eq 2). The product has the reacting
heteroallene incorporated between the metal-hydride bond (M-H). With the hydride now situated on the central carbon of the reacting heteroallene. The exact reactivity can vary depending on the heteroallene used.

\[ \text{LnM}_n - \text{H} \xrightarrow{X=C=X} \text{LnM}_n - X \quad \text{or} \quad \text{LnM}_n / X - C - H \quad (2) \]

\[ X = S, O, NR, CR_2 \]

Organic compounds containing carbonyl functional groups often react in a similar fashion as heteroallenes but the metal binds to the oxygen preferentially in a mono-dentate manner (eq 3).

\[ \text{LnM}_n - \text{H} \quad \xrightarrow{R^1-C=O-R^2} \quad \text{LnM}_n - O - H \quad (3) \]

Protonolysis of the metal hydride involves a reaction with an acid (AH). This reaction releases dihydrogen, resulting in the conjugate base (B) coordinated to the metal centre (eq 4).

\[ \text{LnM}_{n-1} + \text{AH} \quad \xrightarrow{-H_2} \quad \text{LnM}_{n-1} - B \quad (4) \]
1.4 Mercury Compounds

1.4.1 Background

In the elemental form, mercury has been used in various devices including thermometers, float valves and fluorescent lamps. However, the use of elemental mercury has dropped due the concern over its toxicity. Metallic mercury’s electronic configuration strongly resists the removal of an electron and so behaves similarly to a noble gas forming weak intermolecular bonds and thus is a liquid at room temperature (RT).

Mercury(I) is a stable oxidation state and found as the diamagnetic, dimeric cation Hg$_2^{2+}$. Calomel or mercury(I) chloride is a common example and is used as a standard in electrochemistry. Due to mercury’s tendency to bond to itself, mercury(I) will form mercury polycations consisting of linear chains capped with a positive charge. Mercury(II) is the more common oxidation state. Naturally occurring halides, oxides and heavier chalcogens sulfur, selenium and tellurium form stable compounds with mercury(II). Hard-acid-soft-base (HSAB) theory dictates a large soft cation like mercury(II) will have high affinity for a soft anion such as the heavier chalcogens. The mercury chalcogenides and derivatives are semiconductors and have found use as infrared detectors for numerous devices. Mercury has a strong affinity for organic substituents, resulting in stable organo-mercury compounds. These include compounds like methylmercury, which are a dangerous family of compounds that undergo biomethylation and accumulate in waterways, interfering with vital cellular processes of higher organisms upon exposure.

Mercury(II) almost exclusively adopts a linear geometry. The valence electrons are in the 6s orbital. The poorly screened 4f orbitals increase the effective nuclear charge on the valence electrons which in turn increases the Coulombic interaction of the 6s shell to the nucleus. This interaction is absent in zinc and cadmium due to the absence of electrons in the f-orbitals. Relativistic effects and the poorly shielded 6s orbitals provides poor hybridization in the mercury coordination sphere. This results in coordination based on the repulsion of the substituents, thus creating linear coordination.
Mercury has two NMR active nuclei, $^{199}$Hg with a 16.87% natural abundance and $^{201}$Hg with a 13.18% natural abundance. The $^{199}$Hg nuclei are spin 1/2 and produce sharp signals and thus are used for the study of mercury compounds, their structure, dynamics and conformation. $^{199}$Hg coupling to other NMR active nuclei, such as $^1$H, can be observed in an NMR spectrum. The $^{201}$Hg nuclei is quadrupolar and produces broad signals so is not used for characterization and coupling to other NMR active nuclei is not observed.

1.4.2 Coordination Chemistry

β-Diketiminate Mercury Complexes

There have been a limited number of investigations into the coordination chemistry of mercury due to its inherent toxicity. Use of a β-diketiminate ligand as an appropriate ancillary ligand for mercury has been restricted due to the traditional methods of synthesizing group 12 BDI complexes. The general synthesis involves short chained organometallics such as diethyl and dimethyl zinc and cadmium, all of which are toxic and hazardous. Translating this to mercury, organo-mercury complexes would be used where dimethyl mercury is notoriously deadly and much less reactive.$^{26}$

Routes via a more viable reagent such as a metal halide salt have been successful for zinc$^{27}$ and have only been partially successful for cadmium and mercury. An incomplete salt metathesis complex with mercury diiodide and a fluorinated BDI derivative showed that in ethereal solvents, solvent coordination to lithium prevented complete conversion to a β-diketiminato-mercury iodide (I, eq 5).$^{28}$ These complexes are highly soluble, which restricts growth of crystals for structural characterization, and are also very stable, resisting further reactivity.

\[
\begin{align*}
\text{Et}_2\text{O, -40 °C} & \quad \text{HgCl}_2 \\
\text{Li} & \quad \text{N} \quad \text{N} \\
\text{F} & \quad \text{F} \\
\text{F} & \quad \text{F} \\
\text{N} & \quad \text{N} \\
\text{F} & \quad \text{F} \\
\text{O} & \quad \text{Hg} \\
\end{align*}
\]
Earlier attempts at generating a β-diketiminato-mercury complex resulted in a homolytic bound complex (II). Two BDI ligands were bound to the mercury centre through the γ-carbon, which was in equilibrium in solution where one of the ligands switched to the N,N'-binding mode (Scheme 1). Crystallographic data was only obtained for II and showed linear mercury coordination where the equilibrium companion was only observed via $^1$H NMR spectroscopy. This study showed that it was possible for mercury to bind in the N,N'-binding mode of BDI and crystallographic data could be obtained.

![Scheme 1: Reaction and equilibrium of II.](image)

Other related syntheses involved a sila-β-diketiminate derivative where the γ-carbon was replaced by a silicon atom (III, eq 6). Treatment of mercury dichloride in THF with III resulted in a complex with the mercury metal centre bound in a homolytic complex through the silicon of each ligand (IV). This showed that the homolytic complex was favoured when the γ-position was not sterically encumbered. Considering the γ-position is susceptible to electrophilic attack$^{31-32}$ and the high affinity molecular mercury has for carbon atoms, homolytic binding was not unexpected with the silicon derivative III. For a β-diketiminato-mercury that has a terminal substituent for further reactivity, the γ-position of BDI must be blocked to restrict homolytic bonding.

![Equation 6](image)
Mercury Hydrides

Simple mercury hydrides have mostly been computationally studied\textsuperscript{33} with the only mercury hydride on the Cambridge Crystallographic Data Centre (CCDC) being a cluster complex, (CO)\textsubscript{42}Os\textsubscript{18}HgH\textsubscript{34}. This complex consisted of a mercury bridged Os\textsubscript{9} cluster where the hydride ligand was assigned based on indirect methods to be on the mercury centre. This provided little indication on the nature of the mercury hydride or its reactivity.

The terphenyl ligand (terp) has provided enough steric bulk and electronic stabilization to produce group 12 hydride examples.\textsuperscript{35} The mercury hydride derivative (V, eq 7) has been synthesized via a salt metathesis reaction and characterized by \textsuperscript{1}H and \textsuperscript{199}Hg NMR spectroscopy, but the solid state structure was not reported. The \textsuperscript{199}Hg NMR spectrum showed Hg-H coupling with a coupling constant of 2,910 Hz. A resonance, corresponding to the mercury hydride was found at 14.59 ppm in the \textsuperscript{1}H NMR spectrum, but coupling to \textsuperscript{199}Hg was not observed. Computational analysis indicated that a bridged hydride would not be stable in contrast to the iso-structural cadmium and zinc analogues which both form bridged dimers in the solid state. Complex V was only studied in solution, crystallographic comparisons and investigations with other hydrides therefore have not been conducted.

\[
\begin{align*}
\text{Ar} & \quad \text{Hg-I} & \quad \text{KH} & \quad \text{THF} & \quad \text{Ar} \\
\text{Ar} & \quad \text{Hg-H} & & & \quad \text{Ar}
\end{align*}
\] (7)

Organo-mercury hydride compounds are highly unstable and often oligomerize, however this has only been detected with mass spectrometry (MS).\textsuperscript{36} A stable mercury hydride complex (VI) was synthesized by addition of sodium borohydride to a chloride precursor (eq 8). A resonance was found at 13.14 ppm in the \textsuperscript{1}H NMR spectrum of VI, but coupling to \textsuperscript{199}Hg was not observed. The stability of VI was due to the electron-withdrawing effects of the acrylic acid moiety which shortened and stabilized the Hg-H bond.
Thermolysis of mercury hydride VI generated a divinyl-mercury complex, which was postulated to form via a relatively long lived mercury radical intermediate, observed by $^1$H NMR spectroscopy (Scheme 2). This species formed from the net loss of dihydrogen from VI.$^{36}$

Scheme 2: Reported thermolysis and trapping of an organo-mercury radical from VI.

Homonuclear Mercury-Mercury Dimers

In contrast to zinc and cadmium which adopt primarily divalent oxidation states, monovalent mercury as the [Hg$_2$]$^{2+}$ cation, are reasonably common in the literature. Mercury’s affinity to bond to itself provides examples of complexes with oligomeric structures which are often salt-like ionic species.$^{37}$

The first molecular Hg-Hg complexes with alkyl groups were short lived intermediates that were only characterized by NMR spectroscopy and MS. The first dimeric Hg-Hg complex that was structurally characterized was VII (eq 9) revealing a linear molecular R-Hg-Hg-R core with a Hg-Hg bond length of 2.656 Å.$^{38}$ This bond length was longer than the cationic [Hg$_2$]$^{2+}$ and other mercury halide salts with the Hg-Hg distance ranging between 2.49-2.59 Å.

\[ \text{Me}_3\text{Si} - \text{Si} - \text{Si} - \text{H} + \text{^tBu-Hg-Bu} \rightarrow \text{Me}_3\text{Si} - \text{Si} - \text{Hg} - \text{Hg} \]
The precursor used to generate mercury hydride $V$ (eq 7) yielded the homonuclear Hg-Hg dimer ($\text{VIII}$) under reducing conditions (eq 10). The Hg-Hg bond distance in $\text{VIII}$ of 2.574 Å was shorter than the sum of Pauling’s single-bond metallic radii for mercury of 2.88 Å, indicating a bonding interaction was present.

![Equation 10](image)

A molecular mercury-mercury complex bearing a bulky amide ligand on both metal centres was recently synthesized ($\text{IX}$ in Scheme 3). The Hg-Hg bond length in $\text{IX}$ was 2.359 Å. Other structurally characterized relevant complexes bearing Hg-Hg bonds include the collinear tetranuclear R-Co-Hg-Hg-Co-R species, although the Hg-Hg bond length of 2.651 Å was elongated compared to other $[\text{Hg}_2]^{2+}$ cations. Mercury(I) trifluoroacetate exhibited such small crystal dimensions (0.02 x 0.02 x 0.01 nm) that it was only crystallographically characterized explored on a synchrotron. The Hg-Hg bond length in mercury(I) trifluoroacetate was 2.520 Å.

![Scheme 3](image)
1.5 Cadmium Compounds

1.5.1 Background

Like mercury, elemental cadmium has a low melting point but similar to zinc, cadmium will almost exclusively adopt a +2 oxidation state in complexes. However, differing to zinc, cadmium is not biologically relevant but its compounds have found use as coloured pigments, corrosive resistant platings, batteries and semiconductors. Due to its toxicity, cadmium’s use has decreased in battery technologies in the push for more reliable and recyclable elements, and cadmium based pigments and coatings have also been replaced by less toxic materials. Recently, cadmium has been used for semiconductor and solar cell research.43 Cadmium chalcogenide nanoparticles exhibit band gaps dictated by tuneable properties like size and shape, making them desirable for devices requiring specific wavelengths and energies. This is readily observed with quantum dots like CdSe, whose fluorescence energy depends on the size of the CdSe quantum dot.44-45

Similar to mercury, cadmium(I) can be produced as the [Cd₂]^{2+} cation as seen with [Cd₂Cl₆]^{4-}, but these compounds are highly reactive. Most other complexes involve cadmium(II), with a variety of natural oxides, halides and nitrates known. Structures with nucleobases, amino acids and vitamins have also been determined.46 Cadmium complexes share similar coordination geometries with zinc complexes, adopting a tetrahedral coordination when possible. Although not shown to be biologically important, known coordination chemistry indicates a biological affinity and similarity to zinc.

Cadmium has two NMR active nuclei, $^{111}$Cd with a 12.80% natural abundance, and $^{113}$Cd, with a 12.22% natural abundance. Both $^{111}$Cd and $^{113}$Cd are spin 1/2 and can be used to study cadmium containing compounds, although $^{113}$Cd is slightly more sensitive and is the preferred nucleus for cadmium NMR spectroscopy. $^{111}$Cd and $^{113}$Cd coupling to other NMR active nuclei, such as $^1$H, can both be observed in an NMR spectrum.
1.5.2 Coordination Chemistry

β-Diketiminate Cadmium Complexes

The synthesis of β-diketiminato-zinc compounds use the readily available reagent diethyl zinc,\textsuperscript{10} the cadmium alternative is dimethyl cadmium which is less available and significantly more toxic. Two β-diketiminato-cadmium methyl complexes were synthesized by treatment of a protonated β-diketiminate ligand with dimethyl cadmium (X and XII, Scheme 4).\textsuperscript{47} Complex X was used to synthesize another organo-cadmium complex, XI, by treatment with tris-tetrafluorobenzene borane in benzene. X-ray diffraction data showed the cadmium centre was in the plane of the β-diketiminate ligand with an overall distorted trigonal planar geometry for all 3 complexes. When crystals of XI were grown in dimethyl formamide (DMF), a four coordinate geometry at the cadmium centre with a coordinated DMF was found.

As an alternative, cadmium halide salts can also be used to synthesize β-diketiminato-cadmium complexes. Cadmium halide salts are only sparingly soluble in ethereal solvents such as THF and Et\textsubscript{2}O. This results in an incomplete salt metathesis complex when a salt metathesis is attempted with a cadmium halide salt. For instance, treatment of cadmium diiodide with a lithium salt of a BDI ligand in Et\textsubscript{2}O resulted in the formation of bimetallic complexes XIII and XIV (eq 11).\textsuperscript{28} The crystal structure of XIII showed the cadmium centre in a distorted tetrahedral geometry lying in a 6 membered
A β-diketiminato-cadmium complex with a terminal chloride (XVI) had been recently isolated (Scheme 5). The problems with both the use of toxic dimethyl cadmium and the incomplete salt metathesis issue was circumvented by the use of the bis-hexamethyldisilazane cadmium precursor, [Cd(HMDS)_2]. Addition of the β-diketiminate ligand, [BDI_{Dipp-H}], to [Cd(HMDS)_2] resulted in the formation of a β-diketiminato-cadmium amido complex (XV). This was converted to β-diketiminato-cadmium chloride (XVI) by treatment with trimethylsilyl chloride at 90 °C in toluene for 10 days. Complex XVI existed as a dimer in the solid state. Each cadmium centre had a distorted tetrahedral geometry forming a four membered metallocycle through the two bridging chlorides and a six membered metallocycle with the BDI ligand.

Scheme 5: Synthesis of XVI via XV.
Cadmium Hydrides

The bulky trispyrazol borate ligand \([HB(3\text{-}Bu\text{pz})_3]\), or \(\text{Tp}^{\text{Bu}}\), was utilized in the synthesis of the first homonuclear molecular cadmium hydride complex.\(^{51}\) \([\text{Tp}^{\text{Bu}}\text{CdH}]\) (XVII) was synthesized by treatment of cadmium dichloride with a hydride source followed by addition of \([\text{TiTp}^{\text{Bu}}]\) (Scheme 6). The \(^1\text{H}\) NMR resonance for the hydride on complex XVII was at 6.30 ppm, \(^{113}\text{Cd}\) and \(^{111}\text{Cd}\) (\(J_{\text{CdH}}\)) coupling was observed to confirm the presence of a Cd-H bond. A different product was synthesized using a less sterically encumbered trispyrazol borate ligand which contained methyl substituents at the 3 and 5 positions of the pyrazole ring (\(\text{Tp}^{3,5\text{-Me}}\)). A bridging borohydride complex with \(\eta^2\)-coordination to cadmium through two hydrogen atoms (XVIII) generated when KBH\(_4\) was used as the \(\text{H}^-\) source.

![Scheme 6: Synthesis of XVII and XVIII.](image)

Similar to the mercury hydride \(\text{V}\), the terphenyl cadmium hydride (XIX) was generated by addition of KH or NaH to terphenyl cadmium iodide (Scheme 7). Complex XIX was structurally determined as a loosely associated bridged dimer with Cd-H bond distances of 1.786 Å and 2.276 Å. A broad \(^1\text{H}\) NMR resonance at 6.84 ppm was observed with no apparent \(^{113}\text{Cd}\) and \(^{111}\text{Cd}\) coupling. The crystal structure showed distorted trigonal planar conformation at the cadmium centre due to a weak interaction of a neighbouring bridging complex in the unit cell.\(^{35}\) A monomeric variation (XX) was synthesized using 2,4,6-triisopropylphenyl (TIPP) as the flanking aryl group instead of 2,6-diisopropylphenyl (Dipp) (Scheme 7). Structural characterization determined a Cd-H bond distance of 1.794 Å and a similar \(^1\text{H}\) NMR resonance of 6.79 ppm to XIX. For the terminal hydride complex XX the crystal structure showed linear coordination at the cadmium centre.\(^{52}\) Degradation was
not readily observed at room temperature, showing that the increased ancillary ligand bulk was important in the stability of the terminal cadmium hydride.

Scheme 7: Synthesis of XIX and XX.

Complexes XVII and XX have been described as the only organo-cadmium examples where the hydride is at a terminal position making it available for further reactivity. The only known reactivity of the cadmium hydride functionality is the reduction to a homonuclear cadmium-cadmium dimer.\textsuperscript{35}

Homonuclear Cadmium-Cadmium Dimers

The first homonuclear cadmium-cadmium (Cd-Cd) dimer complex (XXI) was synthesized alongside the previously discussed first cadmium hydride (XVII).\textsuperscript{51} The large bulky Tp ligand with methyl substituents at the 3 and 5 positions was used to generate the homonuclear Cd-Cd dimer in a similar fashion as the cadmium hydride (eq 12). By altering the sterics of the ancillary ligand a different cadmium complex was generated. Complex XXI was not structurally determined but was confirmed via \textsuperscript{113}Cd NMR spectroscopy with a \textsuperscript{111}Cd-\textsuperscript{113}Cd coupling constant of 20,646 Hz which indicated a strong bond between the cadmium atoms. Mercury equivalents of these compounds are also known but are less stable than their cadmium counterparts.\textsuperscript{53}
The terphenyl ligand has also been used to generate a homonuclear Cd-Cd dimer derivative (XXII).\textsuperscript{35} Under similar conditions for the synthesis of hydride complexes XIX and XX, but using half the equivalence of sodium hydride, complex XXII was generated (eq 13). The Cd-Cd bond had a bond length of 2.574 Å. In the \textsuperscript{113}Cd NMR spectrum, the \textsuperscript{111}Cd-\textsuperscript{113}Cd coupling constant of 8,650 Hz was smaller than XXI.

\[
\text{Ar}\stackrel{\text{Cd}-\text{Cd}}{\text{Ar}} \text{NaH} \xrightarrow{\text{THF}} \text{Ar}\stackrel{\text{Cd-Cd}}{\text{Ar}}
\]

(13)

A recent molecular Cd-Cd complex bearing a bulky amide ligand on both metal centres was recently synthesized, similar to the mercury analogue IX (XXIII, Scheme 8).\textsuperscript{40} The steric bulk in XXIII prevents further reduction to elemental cadmium. In the \textsuperscript{113}Cd NMR spectrum, the \textsuperscript{111}Cd-\textsuperscript{113}Cd coupling constant was 18,900 Hz, more than double that of XXII, suggesting a stronger and higher s-character Cd-Cd bond than complex XXII. The bond length of XXIII was similar to complex XXII at 2.579 Å.

\[
\begin{align*}
\text{CdI}_2 & \xrightarrow{\text{LK(EntO)}} \text{Ld} \xrightarrow{0.5 (BDIMesMg)_2} \text{Ld-CdL} \\
\text{XXIII} & \quad \text{L} = \text{Ph} \quad R = '\text{Pr}, R' = \text{Me} \\
& \quad \text{R} = \text{Me}, R' = '\text{Pr}
\end{align*}
\]

Scheme 8: Synthesis of XXIII.

The rare examples of homonuclear Cd-Cd examples show that their syntheses requires reducing reagents that can often be too harsh for the ancillary ligand or not harsh enough to reduce the cadmium centre.
1.6 Zinc Compounds

1.6.1 Background

Zinc is the most abundant and available group 12 element.\textsuperscript{3, 16} Several human enzymes and biochemical processes require a zinc metal centre which is obtained through a standard diet.\textsuperscript{54} Zinc is not a typical transition metal since it’s d-orbitals are full and core-like which results in zinc compounds lacking colour.\textsuperscript{55} Zinc is considered a chalcophile due to its affinity for heavier group 11-16 elements and is usually found naturally as ZnS.\textsuperscript{56} As a strongly corrosive resistant metal, the industrial applications of zinc metal include galvanization and alloys, such as brass. Zinc metal is a strong reducing agent and burns in air with a bright blue-green flame.

The +2 oxidation state dominates the chemistry of zinc; however, there are a few examples of zinc in the +1 oxidation state, often containing the highly reactive [Zn\textsubscript{2}]\textsuperscript{2+} core similar to cadmium and mercury.\textsuperscript{57} Zinc complexes are commonly 4 or 6 coordinate, and unlike standard transition metals, zinc bonds have great covalency and stability with nitrogen and sulfur containing substituents.\textsuperscript{56} The zinc chalcogenides are used in various electronic and optic applications.\textsuperscript{58-60} Zinc containing compounds are used as reagents in synthetic chemistry and are heavily researched as catalysts for various organic syntheses.\textsuperscript{61-65}

Zinc has one NMR active nuclei, \textsuperscript{67}Zn with a 4.10% natural abundance which has low sensitivity and produces broad signals with its spin of 5/2. The broad nature of the signal increases with larger compounds, therefore \textsuperscript{67}Zn NMR spectroscopy is not a standard technique and is only used for studying very small complexes of zinc such as zinc dichloride, zinc dibromide and zinc diiodide.
1.6.2 Coordination Chemistry

β-Diketiminate Zinc Complexes

There are many reported β-diketiminato-zinc complexes with varying substituents along the BDI backbone (R₁-R₅, Figure 1). The most commonly used BDI backbones involve R² and R⁴ being methyl groups and R³ or the γ-position as a hydrogen atom. The R¹ and R⁵ positions of the β-diketiminate ligand often differ when used for catalysis. Variations across the backbone include integrated aromatic ring systems providing differing electronic effects.

Several different zinc precursors can be used to generate heteroleptic β-diketiminato-zinc complexes. For instance, organo-zinc precursors, such as dimethyl zinc and diethyl zinc, can react directly with a protonated BDI ligand to form zinc complexes bearing a methyl or ethyl substituent although tert-butyl, phenyl and penta-flurophenyl complexes have also been reported. Zinc bis-amido complexes can also react directly with the protonated BDI ligand to form complexes bearing amide functionality such as hexamethyldisilazane (HMDS). Other common β-diketiminato-zinc amide complexes include a dimethyl amide and diisopropylamide. Zinc halides can also be used, these react with the lithium salt of the BDI ligand to form bridging and terminal β-diketiminato-zinc complexes. Other functionalities include various alkoxides, hydrides, homonuclear Zn-Zn bonds, bis-β-diketiminate complexes and bimetallic species.

Although β-diketiminato-zinc has a vast library of variations, there are still gaps that could further expand their capabilities as reagents or catalysts. Functionalities that are not heavily explored are phosphorus containing substituents, amide derivatives, aryl groups and chalcogenides. Amides and chalcogenides are of particular interest considering zinc’s affinity for these atoms compared to other transition metals and their use in biochemistry.
Zinc Phosphanide Complexes

Most zinc complexes with a phosphorus bond are either involved in clusters, oligomers, are internally bound or have multiple and varying phosphorus atoms within the structure. Although several zinc phosphine complexes are known where the phosphine plays the role of a dative ancillary ligand, relatively few examples of zinc phosphanide complexes are known.

A phosphane, or phosphine, is a derivative of PH$_3$ with organic substituents where a phosphanide ligand is typically an anionic PR$_2$ ligand in which R can be hydrogen, alkyl, aryl or another heteroatom. A phosphide refers to P$^3-$. Basic zinc phosphide (Zn$_3$P$_2$) is a semiconductor with a band gap of 1.5 eV and may have applications in photovoltaic cells. Terminal and accessible zinc phosphanide bonds are rare with only a few examples. The advantage to having a phosphanide ligand is that they are highly sensitive to chemical environments and are easily detectable with $^{31}$P NMR spectroscopy which has a high spectral range from 430 to -250 Hz and a 1/2 spin. Functionally accessible zinc phosphanides could be used as scaffolding for further reactivity with the option of oxidation at the phosphorus centre. To date there is only one β-diketiminato-zinc phosphanide.
complex reported, which was synthesized by treatment of 5 equivalents of phenylphosphine to a β-diketiminato-zinc diethyl complex. This compound was not successfully purified or structurally characterized and was only observed in the $^{31}$P NMR spectrum as a peak at δ -152 ppm with a corresponding doublet in the $^1$H NMR spectrum at δ 4.92 ppm.

Zinc Chalcogenides

There are several zinc complexes, as $[\text{LnZnE-R}]$ (E = S, Se, Te), where thiolates, selenates and tellurates have been bond directly to the metal. Examples mostly involve clusters, oligomers, ligand integrated atoms and mixtures of these. There are few examples involving chalcogenides, referring to the anionic forms, sulfide, selenide and telluride ($E^2$) which maintain a single bond to the metal centre, as terminal substituents. Synthetic routes to these compounds involve many precursors, making their synthetic strategies low on step and atom economy. Zinc sulfide, selenide and telluride nanoparticles are semiconductors with large and controllable band gaps allowing them to be used in various optical and electronic applications including solar cells. Considering the semiconductor nature of zinc chalcogenides it is important to research potential single source precursors.

Only a few examples of β-diketiminato-zinc complexes with chalcogen containing ligands exist. β-Diketiminato-zinc selenide and telluride containing complexes ($XXIV$ and $XXV$) were synthesized by treating a homonuclear Zn-Zn dimer complex with diphenyl diselenide and ditelluride ($\text{Ph}_2E_2$) to produce bridging complexes (Scheme 9). Only the selenide $XXIV$ was structurally characterized and showed a Zn-Se bond distances of 2.429 and 2.595 Å with
E-Zn-E angles close to 90° at 90.99°. Insertion of oxygen into the Te-Zn bond of the telluride \textbf{XXV} was achieved upon addition of O$_2$, the product was structurally characterized and showed the insertion of 2 oxygen molecules.

\begin{figure}
\centering
\includegraphics[width=\textwidth]{scheme9.png}
\caption{Scheme 9: Synthesis of \textbf{XXIV}. Synthesis and reactivity of \textbf{XXV} with O$_2$.}
\end{figure}

Reacting a β-diketiminato-zinc chloride with thallium thiol reagents (Tl-SR) produced bridged dimers \textbf{XXVI} and \textbf{XXVII} (Scheme 10). The Zn-S bond distances of \textbf{XXVI} and \textbf{XXVII} ranged from 2.31-2.42 Å with S-Zn-S bond angles of 90.22° for \textbf{XXVI} and 109.19° for \textbf{XXVII}, the latter contained a bulkier R group. When LiSCPh$_3$ was added, a monomer complex \textbf{XXVIII} was isolated with a Zn-S bond distance of 2.21(4) Å.$^{121}$

\begin{figure}
\centering
\includegraphics[width=\textwidth]{scheme10.png}
\caption{Scheme 10: Synthesis of \textbf{XXVI}, \textbf{XXVII} and \textbf{XXVII}.}
\end{figure}

\textbf{NMR Active Nuclei of Chalcogens}

The NMR spectroscopy of the NMR active nuclei of the chalcogens can be useful characterization tools when investigating complexes containing varying amounts of the chalcogen. The least useful, in terms of NMR characterization, is sulfur due to the low abundance (0.76%) of its only NMR active nuclei, $^{33}$S, which has a 3/2 spin producing broad
signals. The signals are broader with larger and more asymmetric compounds leading to $^{33}$S NMR spectroscopy only being used for the characterization of small sulfur containing compounds such as dimethyl sulfoxide (DMSO). Coupling to other NMR active nuclei is not observed in other NMR spectra due to the low abundance of the $^{33}$S nuclei.

Selenium has one NMR active nuclei, $^{77}$Se with a 7.63% natural abundance and a spin of 1/2, producing narrow signals. The $^{77}$Se nuclei can couple to other NMR active nuclei, such as $^{31}$P, which can be observed and used to characterize the selenium nuclei environment. Tellurium has two NMR active nuclei, $^{125}$Te with a 7.07% natural abundance and $^{123}$Te with a 0.89% natural abundance. Both nuclei are 1/2 spin and produce narrow signals but $^{125}$Te is used for tellurium NMR since $^{125}$Te is more abundant, has higher sensitivity and yields sharper signals. The $^{125}$Te nuclei can couple to other NMR active nuclei, such as $^{31}$P, which can be observed and used to characterize the tellurium nuclei environment.

1.6.1 Research from 2014

Research with novel zinc coordination was conducted as part of a Summer Research Project in 2014. This work is related to zinc phosphanides, zinc chalcogenides and later work in Chapter 2 of this thesis and so has been included as part of this introduction.

The use of zinc halide salts is problematic since they are hygroscopic. The presence of water can produce undesired protonation products when these halide salts are used for the synthesis of water sensitive compounds. The standard purification method for zinc dichloride is to recrystallize zinc dichloride from hot dioxane. This method proved unsuccessful in producing water free zinc dichloride and so zinc dibromide was considered. The standard purification method for zinc dibromide is sublimation. With the correct sublimation gear and heating apparatus it was possible to sublime zinc dibromide under vacuum at temperatures above 350 °C. The purified zinc dibromide was used in a known synthesis to generate the starting material [BDIDippZnBr] (XXIX). The common BDI ligand, BDIDipp, was used which contains two Dipp groups at R1 and R5, methyl groups at R2 and R4 and a hydrogen atom known as the γ-proton at R3 (Figure 1).
Synthesis of Zinc Phosphanide Complexes

\[
\begin{align*}
\text{Ar} & \quad \text{Zn} \quad \text{Ar} \\
\text{Br} & \quad \text{Li-PR}_2 \\
\text{Et}_2\text{O}, 16 \text{ hr} & \quad \text{R} = \text{Cy (XXX), Ph (XXXI)} \\
\end{align*}
\]

Attempts to generate zinc-phosphanide complexes from lithium phosphanide salts were attempted via a salt metathesis reaction with **XXIX**. The lithium phosphanide salts, lithium dicyclohexylphosphane (Li-PCy₂) and lithium diphenylphosphane (Li-PPh₂), were generated by the treatment of the corresponding phosphine with \(n\)-butyllithium (\(n\text{BuLi}\)). \(\beta\)-Diketiminato-zinc dicyclohexylphosphane was synthesized by treatment of **XXIX** with Li-PCy₂. To minimize the possibility of incomplete salt metathesis, non-ethereal solvents were initially used. However, no reaction was observed in those solvents. When the reaction was performed in diethyl ether the zinc dicyclohexylphosphane (**XXX**) was produced with no evidence of an incomplete salt metathesis (eq 14).

The \(^1\text{H}\) NMR spectrum of **XXX** shows one septet at \(\delta 3.22\) ppm (formally a quartet of quartets), indicating a single environment for the methine proton on the isopropyl groups and a singlet at \(\delta 5.00\) ppm for the \(\gamma\)-proton, downfield to the \(\gamma\)-proton of the precursor **XXIX**. There is an inclusion of cyclohexyl signals at 1.56 and 1.28 ppm. The \(^{31}\text{P}\) NMR spectrum shows a single peak at \(\delta -22.2\) ppm, slightly downfield to Li-PCy₂ (\(\delta -21.9\) ppm).
Figure 9: ORTEP diagram of XXX showing thermal ellipsoids at 30% probability. Only bonds for P2 are shown. One set of cyclohexyl peaks are shown (a). Hydrogens omitted for clarity.

Yellow cube shaped crystals suitable for an X-ray diffraction study were grown from diethyl ether at -30 °C. Complex XXX is monomeric in the solid state and showed disorder at the phosphorus nucleus and the two cyclohexyl groups. Both positions were modelled and what are shown in Figure 9 are the nuclei with the highest occupancy (81%).

The nitrogens of the BDI ligand do not show any amide or imine character and are statistically indistinguishable from each other, indicative of a delocalized π-system in the BDI backbone. Complex XXX has a BDI bite angle of 95.5(1)°.

The Zn-P2 bond length of 2.285(1) Å is shorter than a dative phosphane bond with bond lengths ranging 2.393-2.401 Å\textsuperscript{93} and a P-metallated phosphasilene with a Zn-P bond length of 2.367 Å.\textsuperscript{97} The Zn-P2 bond length in complex XXX is closer to that of an internally bound zinc phosphane in a cluster with bond lengths ranging from 2.302 to 2.334 Å.\textsuperscript{95} The phosphane ligand is asymmetric to the zinc metal centre, skewed towards N2 as represented with a shorter N2-Zn-P2 bond angle of 123.8(1)° compared to the N1-Zn-P2 bond angle of 137.7(1)°.
Table 1: Selected bond lengths (Å) and angles (°) for XXX.

<table>
<thead>
<tr>
<th>Bond</th>
<th>Length</th>
<th>Bond</th>
<th>Length</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zn-P2</td>
<td>2.285(1)</td>
<td>Zn-N1</td>
<td>1.971(4)</td>
</tr>
<tr>
<td>N1-C1</td>
<td>1.318(6)</td>
<td>C1-C2</td>
<td>1.398(6)</td>
</tr>
<tr>
<td>C3-N2</td>
<td>1.329(6)</td>
<td>N1-C6</td>
<td>1.441(6)</td>
</tr>
<tr>
<td>P2-C30a</td>
<td>1.942(5)</td>
<td>P2-C36a</td>
<td>1.856(5)</td>
</tr>
<tr>
<td>N1-Zn-N2</td>
<td>95.5(1)</td>
<td>Zn-N1-C1</td>
<td>122.4(3)</td>
</tr>
<tr>
<td>C1-C2-C3</td>
<td>127.9(4)</td>
<td>C2-C3-N2</td>
<td>124.2(4)</td>
</tr>
<tr>
<td>N1-Zn-P2</td>
<td>137.7(1)</td>
<td>N2-Zn-P2</td>
<td>123.8(1)</td>
</tr>
<tr>
<td>Zn-P2-C36a</td>
<td>110.1(2)</td>
<td>C30a-P2-C36a</td>
<td>103.8(2)</td>
</tr>
</tbody>
</table>

Addition of Li-PPh₂ to a diethyl ether solution of zinc bromide XXIX resulted in the formation of multiple products. The ¹H NMR spectrum of the reaction mixture revealed [BDI_{Dipp}-Li] and potentially two new BDI containing compounds as indicated by the presence of two signals (δ 5.02 and 5.00 ppm) which typically correspond to the γ-proton of the BDI backbone. The ³¹P NMR spectrum shows a single peak at δ -45.8 ppm, up-field of Li-PPh₂ (δ -43.9 ppm).
Figure 10: ORTEP diagram of XXXII showing thermal ellipsoids at 30% probability. Hydrogens omitted for clarity.

Yellow cube shaped crystals suitable for an X-ray diffraction study were grown from diethyl ether at -30 °C over a week. The product was not the expected zinc phosphanide complex XXXI and was instead an incomplete salt metathesis complex with two diethyl ether molecules coordinated to the lithium centre (XXXII) in a similar manner to other group 12 salt metathesis complexes I and XIII.

Complex XXXII is monomeric in the solid state. The zinc centre lies 0.412 Å out of the plane consisting of N1-C1-C2-C3-N2 (referred to herein as BDI plane). The plane consisting of O1-Li-O2 is 8.74° from the BDI plane. The bromide atoms are not equivalent based on their bond lengths to zinc, 2.4581(4) and 2.4172(4) Å, but their bond lengths only differ slightly to the lithium, 2.549(4) and 2.553(5) Å. The BDI backbone has delocalization as observed by the statistically indistinguishable C-C bond lengths (1.390(3) and 1407(4) Å) and N-C bond lengths (1.329(3) and 1.325(3) Å). Complex XXXII has a smaller bite angle for the BDI ligand compared to the zinc phosphanide XXX due to the accommodation of large bromide atoms. The bite is in range of existing bridged β-diketiminato-zinc bromide complexes which exhibit a BDI bite angle range of 96.67-98.33°.28
The BDI plane is 84.38° from the plane consisting of Zn-Br1-Li-Br2. The metallocycle with lithium is square shaped, the Zn-Br1-Li and Zn-Br2-Li bond angles (82.3(1)° and 83.0(1)°) are shorter than the Br1-Zn-Br2 and Br1-Li-Br2 bond angles (99.33(2)° and 93.5(2)°). The Li-O bond lengths of 1.940(5) and 1.995(5) Å, and the variable bond angles about the lithium centre indicates asymmetry at the lithium metal centre with the Et₂O molecules.

The zinc-bromide bond lengths are longer than standard zinc dibromide at 2.21 Å\(^{124}\) potentially due to the bridging geometry. The Zn-Br bond lengths follow the same trend as a bridged β-diketiminato-zinc bromides which have shorter bond lengths ranging 2.402-2.408 Å for one bromide and longer bond lengths ranging 2.418-2.435 Å for the other bromide.\(^{28}\) Both bromide atoms are formally Br\(^2-\) contributing 0.5- to zinc and 1.5- to lithium. The Br1-Li-Br2 functionality is a bidentate ligand, formally contributing a single charge to Zn(II).

**Table 2:** Selected bond lengths (Å) and angles (°) for XXXII.

| Bond Lengths | Zn-N1 | Zn-N2 | Zn-N1-C1 | Zn-Br1 | Zn-Br2 | Br1-Li | Br2-Li | Li-O1 | Li-O2 | N1-Zn-N2 | Zn-N1-C1 | N1-C1-C2 | N1-Zn-Br1 | N1-Zn-Br2 | O1-Li-O2 | O1-Li-Br2 | O2-Li-Br2 |
|--------------|-------|-------|----------|--------|--------|--------|--------|------|------|----------|----------|----------|----------|----------|----------|----------|----------|----------|
| Å            | 1.975(2) | 1.978(2) | 1.329(3) | 2.4581(4) | 2.4172(4) | 2.549(4) | 2.553(5) | 1.940(5) | 1.995(5) | 97.69(8) | 120.3(2) | 123.6(2) | 83.0(1) | 113.61(6) | 106.6(2) | 104.6(2) |
Unfortunately repeated attempts at obtaining pure diphenylphosphanide XXXI were unsuccessful with only contaminated $^1$H NMR and $^{31}$P NMR spectra indicating diphenylphosphanide XXXI as a possible product. Complex XXXII is potentially the product of a degradation pathway.

Addition of Chalcogens into Zinc Phosphanide Bonds

Addition of selenium powder to a Et$_2$O solution of the dicyclohexylphosphanide XXX resulted in the formation of the dicyclohexylphosphinodiselenoate complex XXXIII (eq 15). The $^1$H NMR spectrum of XXXIII shows one septet at $\delta$ 3.49 ppm, indicating a single environment for the methine proton on the isopropyl groups and a singlet at $\delta$ 4.85 ppm for the $\gamma$-proton, up-field from dicyclohexylphosphanide XXX. The $^{31}$P NMR spectrum shows a single peak at $\delta$ 51.2 ppm, downfield from XXX, with $^{77}$Se satellites. The $^{77}$Se-$^{31}$P coupling constant of 451.5 Hz is consistent with the addition of two selenium atoms to a phosphorus atom seen in $\beta$-diketiminato-tin and germanium phosphanide complexes.$^{125}$ Throughout the reaction no other signals were observed in the $^{31}$P NMR spectrum that would indicate a complex with only one selenium atom added, a dicyclohexylphosphinoselenoite complex.
Orange rectangular crystals suitable for an X-ray diffraction study were grown from diethyl ether at -30 °C. Complex XXXIII is monomeric in the solid state with a molecule of diethyl ether in the unit cell. The phosphorus atom is formally P^5+ and the selenium atoms are both Se^2-. The Zn-Se bond lengths of 2.5002(4) and 2.5010(5) Å are very similar where the Se-P bond lengths of 2.1708(4) and 2.1737(5) Å only differ slightly. In contrast to β-diketiminato-germanium dicyclohexylphosphinodiselenoate complexes, which are three coordinate, the dicyclohexylphosphinodiselenoate ligand in complex XXXIII is bidentate resulting in a tetrahedral zinc centre. The Zn-Se bonds are in range of the Zn-Se bonds in the selenium bridged BDI complex XXIV which range from 2.429-2.595 Å. Other Zn-Se bonds from various compounds range from 2.320-2.609 Å. Attempts to produce a mono-insertion product proved unsuccessful even with treatment of a single equivalent or less of selenium powder.

The nitrogens of the BDI ligand do not show any amide or imine character and are statistically indistinguishable from each other, indicative of a delocalized π-system in the BDI
backbone. Complex **XXXIII** has a slightly larger BDI bite angle of 99.07(5)° to the starting material dicyclohexylphosphaneide **XXX**.

The plane consisting of N1-C1-C2-C3-N2 is 89.61° from the plane of the phosphinodiselenoate ligand. The phosphinodiselenoate ligand is square shaped, the Zn-Se1-P and Zn-Se2-P bond angles (80.25(2) and 80.29(2)°) are shorter than the Se1-Zn-Se2 and Se1-P-Se2 bond angles (90.11(2) and 109.13(2)°).

**Table 3:** Selected bond lengths (Å) and angles (°) for **XXXIII**.

<table>
<thead>
<tr>
<th>Bond</th>
<th>Length/Angle</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zn-N1</td>
<td>1.971(1)</td>
</tr>
<tr>
<td>Zn-N2</td>
<td>1.974(1)</td>
</tr>
<tr>
<td>N1-C1</td>
<td>1.327(2)</td>
</tr>
<tr>
<td>C1-C2</td>
<td>1.405(2)</td>
</tr>
<tr>
<td>N1-C6</td>
<td>1.437(2)</td>
</tr>
<tr>
<td>Zn-Se1</td>
<td>2.5010(5)</td>
</tr>
<tr>
<td>Zn-Se2</td>
<td>2.5002(4)</td>
</tr>
<tr>
<td>P-C30</td>
<td>1.828(1)</td>
</tr>
<tr>
<td>Se1-P</td>
<td>2.1708(4)</td>
</tr>
<tr>
<td>Se2-P</td>
<td>2.1737(5)</td>
</tr>
<tr>
<td>N1-Zn-N2</td>
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<td>C1-C2-C3</td>
<td>130.9(1)</td>
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<td>116.19(4)</td>
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<td>C30-P-Se1</td>
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</tr>
<tr>
<td>C36-P-Se2</td>
<td>110.81(5)</td>
</tr>
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</table>

Addition of selenium powder to a Et₂O solution of zinc diphenylphosphaneide complex **XXXI** resulted in the formation of a diphenylphosphinodiselenoate, complex **XXVI** (eq 15). Although impure diphenylphosphaneide **XXXI** was used, **XXXIV** was isolated as a single compound. The ¹H NMR spectrum of **XXXIV** shows one septet at δ 3.53 ppm, indicating a single environment for the methine proton on the isopropyl groups and a singlet at 4.84 ppm, up-field to the suspected phosphaneide **XXXI** γ-proton signal. The ³¹P NMR spectrum shows a single peak at δ 6.3 ppm with a ⁷⁷Se-³¹P coupling constant of 472.7 Hz, similar to the dicyclohexylphosphinodiselenoate **XXXIII**, indicating two selenium atoms had been added into the zinc-phosphaneide bond. Throughout the reaction no other signals were observed in
the $^{31}$P NMR spectrum that would indicate a complex with only one selenium atom added, a diphenylphosphinoselenoite complex.

Figure 12: ORTEP diagram of XXXIV showing thermal ellipsoids at 30% probability. Hydrogens omitted for clarity.

Orange rectangular crystals suitable for an X-ray diffraction study were grown from diethyl ether at -30 °C. Complex XXXIV is monomeric in the solid state. The phosphorus atom is formally P$^{5+}$ and the selenium atoms are both Se$^{2-}$. The Zn-Se bond lengths, Zn-Se1 of 2.4749(5) and Zn-Se2 of 2.5235(3) Å, differ where the Se-P bond lengths of 2.1588(6) and 2.1671(5) Å are similar. In contrast to β-diketiminato-germanium diphenylphosphinodiselenoate complexes, which are three coordinate, the diphenylphosphinodiselenoate ligand in complex XXXVI is bidentate resulting in a tetrahedral zinc centre. The Zn-Se bonds are in range of the Zn-Se bonds in the selenium bridged BDI complex XXIV which range from 2.429-2.595 Å. Other Zn-Se bonds from various compounds range from 2.320-2.609 Å. Attempts to produce a mono-insertion product proved unsuccessful even with treatment of a single equivalent or less of selenium powder.
The nitrogens of the BDI ligand do not show any amide or imine character and are statistically indistinguishable from each other, indicative of a delocalized π-system in the BDI backbone. Complex XXXVI has a similar BDI bite angle of 99.23(5)° to dicyclohexylphosphinodiselenoate XXXIII.

The plane consisting of N1-C1-C2-C3-N2 is 89.70° from the plane of the phosphinodiselenoate ligand. The phosphinodiselenoate ligand is square shaped, the Zn-Se1-P and Zn-Se2-P bond angles (80.22(2) and 81.18(2)°) are shorter than the Se1-Zn-Se2 and Se1-P-Se2 bond angles (89.47(3) and 108.84(2)°). The phosphinodiselenoate ligand shares similar bond angles to the phosphinodiselenoate ligand of dicyclohexylphosphinodiselenoate XXXIII.

**Table 4:** Selected bond lengths (Å) and angles (°) for XXXIV.

<table>
<thead>
<tr>
<th>Bond Length/Angle</th>
<th>Value</th>
</tr>
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<td>Se2-P-C30</td>
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</table>
Addition of tellurium powder to the dicyclohexylphosphanide XXX did not give the double addition product. Instead, a mono-insertion product was suspected, a dicyclohexylphosphinotellunoite, complex XXXV (eq 16).

The $^1$H NMR spectrum of the reaction mixture has both $[$BDI$_{Dipp}$-H$]$ and a potentially new BDI containing complex as indicated by the presence of a signal ($\delta$ 5.06 ppm) in the $^1$H NMR spectrum which typically corresponds to the $\gamma$-proton of the BDI backbone. The $^{31}$P NMR spectrum shows a peak at $\delta$ 7.2 ppm with a pair of satellites. Only the satellites from $^{125}$Te were observed with a $^{125}$Te-$^{31}$P coupling constant of 285.0 Hz. This is a smaller coupling constant compared to the phosphinodiselenoate complexes XXXIII and XXXIV. This coupling is similar to $\beta$-diketiminato-germanium dicyclohexylphosphinotellunoite complexes, thus we suspect the new BDI containing complex to be the mono-inserted product (XXXV).$^{125}$ Two types of crystals appeared to grow out of an orange reaction mixture at -30 °C, yellow needles and yellow cubes of which could not be separated. Both crystals were not suitable for an X-ray diffraction study.

Attempts to insert sulfur into the zinc-phosphanide bond of the dicyclohexylphosphanide complex XXX to synthesize a dicyclohexylphosphinodithioate proved unsuccessful. Additions of elemental sulfur resulted in multiple products, detected by $^1$H and $^{31}$P NMR spectroscopy, which were suspected to be both the single and double addition products. Isolation and determination of the products in the reaction mixture was inconclusive.
1.7 Pre-Thesis Research Motive

The recent synthesis of the cadmium chloride (XVI) was the basis for some of the following research outlined in Chapter 2. Further investigation into complex XVI as a viable synthon to other novel cadmium complexes was influenced by pre-thesis research (December 2015) on the reduction of the cadmium centre to a cadmium hydride (3). Optimized synthetic details were further investigated for complexes XVI and 3 during this project and have been included in Chapter 2. This created interest in developing a suitable mercury synthon for similar reductions which required the development of a compatible BDI ancillary ligand. The resulting BDI ancillary ligand used for mercury was also used with zinc to act as a comparison and was the basis of further research into novel zinc functionalities.

1.7.1 Synthesis of $[\text{Ar-BDI}_{\text{Dipp}}^{-}\text{H}]$

Due to reported unsuccessful attempts with BDI and mercury, an alternative BDI ligand was used. The alternative BDI had success with copper(II) complexes that were found to form similar homolytic complexes.\textsuperscript{126} Due to the incorporation of an aryl group across the BDI backbone, the ligand is denoted as $\text{Ar-BDI}_{\text{Dipp}}$ where Dipp refers to the flanking 2,6-diisopropylphenyl groups at positions $R^1$ and $R^5$ (Figure 3). Ligand synthesis is according to a previously published procedure (Scheme 11).\textsuperscript{126} Addition of 2'-fluoroacetophenone to titanium tetrachloride (TiCl$_4$) and 2,6-diisopropylaniline in hexane produced a dense brown precipitate. The imine intermediate was isolated by filtration and dissolved in THF, lithium 2,6-diisopropylanilide was added, after which the reaction mixture changed from a dark orange to a dark yellow solution. Following three days under reflux and workup, the ligand, $[\text{Ar-BDI}_{\text{Dipp}}^{-}\text{H}]$, was crystallized as pale yellow-orange crystals from a 1:1 methanol: dichloromethane (DCM) solution.

\begin{equation}
\begin{array}{cccccc}
\text{F} & \text{O} & \text{NH}_2 & + & 3 & \text{Hex. 75 °C, 16 hr} \\
\text{F} & \text{N} & + & \text{THF, 90 °C, 72 hr} \\
\text{Dipp-N(H)Li} & & & & & \\
\end{array}
\end{equation}

\begin{equation}
\text{Scheme 11: Synthesis of $[\text{Ar-BDI}_{\text{Dipp}}^{-}\text{H}]$.}
\end{equation}
Chapter 2

Results and Discussion

2.1 Mercury

2.1.1 Synthesis of β-Diketiminato-mercury Chloride

The standard procedure to create the lithium salt of a BDI ligand is to use nBuLi as a strong base. To prevent residual nBuLi reacting with the mercury halide salt, lithium diisopropylamide (LDA) was used as an alternative strong base as demonstrated in the synthesis of bis-β-diketiminato-mercury II. To a toluene solution of [Ar-BDI\text{dipp}-H], a toluene slurry of LDA was added. The light yellow solution was left stirring overnight, after which a dark yellow precipitate had formed. After removing the volatiles \textit{in vacuo}, the lithium salt, [Ar-BDI\text{dipp}-Li], was isolated as a dark yellow powder. Impurities were further removed by triturations in hexane.

Complex 1 was synthesized by treatment of a toluene slurry of mercury dichloride with a toluene solution of fresh [Ar-BDI\text{dipp}-Li] (Scheme 12). After 24 hours at room temperature the reaction mixture was filtered through celite and the volatiles removed \textit{in vacuo}, yielding a light yellow solid (1).

\[ \text{Scheme 12: Synthesis of [Ar-BDI\text{dipp}-Li] followed by the synthesis of 1.} \]
The $^1$H NMR spectrum of 1 shows two septets at δ 3.48 and 3.11 ppm corresponding to two different environments for the methine proton on the isopropyl groups. Four doublets for the isopropyl methyl groups are found at δ 1.33, 1.24, 1.14 and 1.04 ppm, producing evidence of an asymmetrical ligand environment. The $^1$H NMR resonances differ from that of [Ar-BD]Dipp-Li).

![Image of molecular structure](image)

**Figure 13:** ORTEP diagram of 1 showing thermal ellipsoids at 30% probability. Hydrogens omitted for clarity.

Yellow floret-shaped crystals suitable for an X-ray diffraction study were grown from hexane at -30 °C. Complex 1 is monomeric in the solid state. The mercury centre has a large angle between N2 and Cl with a N2-Hg-Cl angle of 169.18°. The N2-Hg bond length of 2.044(2) Å is significantly shorter than the N1-Hg bond length of 2.444(2) Å, consistent with a more anionic N2-Hg bond and a dative N1-Hg, with the N1-Hg-N2 bond angle approaching 90° (81.33(7)°). The N1-C1 bond length of 1.441(3) Å is shorter than the N2-C3 bond length of 1.282(3) Å, indicating N1 retains its imine character whereas N2 has more amido-imine character. Nitrogens with the amido-imine character are henceforth referred to as an amide. This type of bonding is commonly observed for three coordinate mercury complexes.
where the mercury centre remains relatively linear with an added stabilizing dative bond influencing the linear character of the mercury centre.127-129

The plane consisting of N1-C1-C2-C3-N2 (referred to herein as BDI plane) exhibits little distortion. The mercury atom is displaced by 1.172 Å from the BDI plane due to the unusual distorted trigonal planar geometry of the mercury metal centre. The angle between the BDI plane and the plane consisting of N1-N2-Hg-Cl is 44.21°.

Table 5: Selected bond lengths (Å) and angles (°) for 1.

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<tr>
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<td>Hg-N2</td>
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</tr>
<tr>
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</table>

This is the first structurally characterized 3-coordinate β-diketiminato-mercury complex; unfortunately further derivatization of 1 was unsuccessful. Reactions with lithium salts including Li-PCy2, Li-PPh2 and lithium hexamethyldisilazane ([Li-HMDS]) yielded [Ar-BDI_{dipp}-Li] and small amounts of [Ar-BDI_{dipp}-H]. Attempts to synthesize a hydride using lithium tetraethyl borohydride and potassium hydride proved unsuccessful, yielding similar results. The unique structure, geometry and strong Hg-Cl bond in complex 1 possibly hindered salt metathesis with lithium salts and hydride sources. A more pliable substituent, such as HMDS, may yield better reactivity.
2.2 Cadmium

2.2.1 Alternative Synthesis of β-Diketiminato-cadmium Chloride

A new route to β-diketiminato-cadmium chloride (XVI) was used in this project. This new route involves two steps instead of three and allows the synthesis of the chloride XVI in three days instead of ten (Scheme 13).\(^{49}\) The intermediate, [HMDS-CdCl], was synthesized \textit{in situ} by addition of one equivalent of [Li-HMDS] to a THF slurry of cadmium chloride. After 24 hours at room temperature, the volatiles were removed \textit{in vacuo} to yield a thick grey oil that contained a single peak in the \(^1\)H NMR spectrum at \(\delta 0.289\) ppm, downfield to both [Li-HMDS] (\(\delta 0.135\) ppm) and [CdHMDS\(_2\)] (\(\delta 0.210\) ppm). Cadmium chloride XVI was synthesized by filtering a toluene solution of [HMDS-CdCl], adding a toluene solution of [BDI\(_{\text{Dipp}}\)-H] and stirring for 72 hours at 55 °C. Volatiles were removed \textit{in vacuo}, triturations with hexane further removed impurities. Pure cadmium chloride XVI was obtained by crystallization from toluene.

\[
\text{CdCl}_2 + [\text{Li-HMDS}] \xrightarrow{\text{THF, RT, 12 hrs}} [\text{HMDS-CdCl}] \xrightarrow{[\text{BDI}_{\text{Dipp}}\text{-H}] \text{Tol. 55 °C, 72 hrs}} \text{XVI}
\]

\textbf{Scheme 13: Alternative synthesis of XVI.}

2.2.2 Derivatization of β-Diketiminato-cadmium Chloride

Phosphanide Synthesis

Attempts to generate cadmium-phosphanide complexes from lithium phosphanide salts were attempted via a salt metathesis reaction with XVI. The lithium phosphanide salts, Li-PC\(_2\) and Li-PPh\(_2\), were used. To remove the possibility of an incomplete salt metathesis in the reaction with chloride XVI, a non-coordinating solvent, toluene, was used.

Cadmium dicyclohexylphosphanide (2) was synthesized by addition of cadmium chloride XVI to a toluene slurry of Li-PC\(_2\). After 16 hours the resulting reaction mixture was filtered
through celite and the volatiles were removed in vacuo yielding the sticky yellow residue of complex 2 (eq 17). Treatment of cadmium chloride XVI with Li-PPh₂ led to multiple products.

The \(^1\)H NMR spectrum of the reaction mixture is not consistent with a single compound and shows [BDI\textsubscript{Dipp}-Li] and [BDI\textsubscript{Dipp}-H] alongside the suspected new product (phosphanide 2) indicated by a new singlet from the γ-proton at δ 4.92 ppm, downfield to the γ-proton of chloride XVI (δ 4.64 ppm). The \(^{31}\)P NMR spectrum shows a single peak at δ 5.4 ppm with \(^{113}\)Cd and \(^{111}\)Cd satellites which have coupling constants of 570.8 Hz and 545.8 Hz respectively.

The coupling between cadmium and phosphorus can be related to existing compounds. The coupling \(^1\)J\textsubscript{PCd}, seen in \(^{31}\)P NMR spectroscopy, is around 1500 Hz for a phosphine\textsuperscript{130-131} and around 700 Hz for a phosphanide.\textsuperscript{132} Although complex 2 has a more up field shift to existing cadmium phosphanides, it was determined that the phosphanide had been successfully bound to the cadmium centre in complex 2 from a salt metathesis reaction. Attempts to produce a crystalline product suitable for an X-ray diffraction study proved unsuccessful. The reaction always produced multiple degradation products which would increase in abundancy over time, as such; we were unable to grow crystals of complex 2.

**Hydride Synthesis**

Several hydride sources were screened in order to produce a cadmium hydride from chloride XVI. This included potassium hydride, sodium hydride and calcium hydride of which only lithium triethylborohydride, or Super Hydride, resulted in the clean formation of the desired cadmium hydride (3). Complex 3 was synthesized by the slow addition of Super Hydride to a toluene or diethyl ether solution of chloride XVI (eq 18). The resulting residue
was re-dissolved in a minimum amount of hexane or diethyl ether and kept at -30 °C to produce 3 as a crystalline solid.

\[
\text{Cl} \quad \text{Li-BE} \text{t}_3 \text{H} \quad \text{Tol. or Et}_2 \text{O 30 min.}
\]

The $^1$H NMR spectrum of 3 shows one septet at δ 3.29 ppm, indicating a single environment for the central proton on the isopropyl groups, and a singlet from the γ-proton at δ 4.91 ppm, downfield to that of chloride XVI. The hydride resonance was found as a very broad signal at δ 5.63 ppm which was upfield to reported cadmium hydride complexes.\(^{35}\) Due to the broad nature of the hydride peak, cadmium satellites were not observed. The broad signal could be due to coupling to the quadrupolar $^{14}$N nuclei.

Figure 14: ORTEP diagram of 3 showing thermal ellipsoids at 30% probability. Selected hydrogens omitted for clarity.

Small white cubic crystals suitable for an X-ray diffraction study were grown from hexane at -30 °C. Complex 3 is monomeric in the solid state. The 3-coordinate cadmium centre is
co-planar with the BDI backbone. The location of the hydride was calculated from residual
densities in the difference density map.

The N1-Cd and N2-Cd bond lengths 2.160(1) and 2.163(1) Å are statistically indistinguishable
from each other and show symmetrical bonding to both nitrogens of the ligand. The bite
angle, N1-Cd-N2, of 88.45(5)° is similar to other three coordinate β-diketiminato-cadmium
complexes involving Dipp groups at the R¹ and R⁵ positions, BDI organo-cadmium complexes
X and XI, with bite angles of 88.45° and 90.54°, respectively. The bite angle is larger than
Ar-BDI organo-cadmium XII which used a variation of the ligand with a smaller bite angle
range of 86.79-87.35°. The N1-C₁ and N2-C₃ bond lengths of 1.332(2) and 1.328(2) Å and
N1-C₆ and N2-C₁₈ bond lengths of 1.432(2) and 1.430(2) Å are statistically indistinguishable
from each other, indicative of a delocalized π-system in the BDI backbone.

Table 6: Selected bond lengths (Å) and angles (°) for 3.

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<th>Lengths (Å)</th>
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<td>Cd-N2</td>
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<tr>
<td>N1-C6</td>
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<td>N2-C₁₈</td>
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<td>C3-N2-Cd</td>
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</table>

The hydride in complex 3 is comparable to the terphenyl hydride complex XX’s as they are
both terminal cadmium hydride complexes. Similar Cd-H stretches are found, 1734 cm⁻¹ for
3 and 1735 cm⁻¹ for monomeric terphenyl cadmium hydride XX. Density functional theory
(DFT) calculations with THF and 3 showed the cadmium centre does not possess the
geometry or available orbitals required to expand its coordination, seen with bridging
terphenyl cadmium hydride XIX. This was confirmed experimentally as addition of THF does
not alter the chemical shifts in the ¹H NMR spectrum at room temperature or with increased
temperatures. A melting point of hydride 3 was not obtained as it degrades to a black
material upon heating. Slow degradation of hydride 3 was observed in the ¹H NMR spectrum with [BDIDipp-H], small amounts of a homonuclear Cd-Cd dimer (5) and a black precipitate, accumulating after two weeks.
2.2.3 Reactivity of β-Diketiminato-cadmium Hydride

Reactivity with Carbodiimides

The nucleophilicity of the cadmium hydride was examined by the reaction of complex 3 with dicyclohexylcarbodiimide (DCC), diisopropylcarbodiimide (DIC) and ditertbutylcarbodiimide (DBC). Similar reactivity studies have been conducted with β-diketiminato-tin and β-diketiminato-zinc hydrides to give a net insertion of the carbodiimide into the metal hydride bond. As such, with this cadmium hydride an insertion of the carbodiimide was expected to produce the cadmium amidinate, complex 4 (eq 19).

\[ \text{Ar}_2\text{N}^+\text{Cd} \quad \text{H} \quad \text{R-N=C=N-R} \quad \text{Hex. 16 hr} \]

\[ \text{R} = \text{Cy, } \text{iPr, } \text{tBu} \]

However when DIC or DCC was added to hydride 3, the product was not the expected insertion amidinate product 4 but was instead the homonuclear Cd-Cd dimer (5).

\[ 2\text{Ar}_2\text{N}^+\text{Cd} \quad \text{H} \quad \text{R-N=C=N-R} \quad \text{Hex. 16 hr} \]

\[ \text{R} = \text{Cy, } \text{iPr} \]

(20)

The \(^1\text{H} \) NMR spectrum of 5 shows one septet at δ 3.12 ppm, indicating a single environment for the central proton on the isopropyl groups and a singlet at δ 4.78 ppm, up-field to hydride 3. When the reaction was monitored by \(^1\text{H} \) NMR spectroscopy, \( \text{H}_2 \) was observed in the reaction mixture.
Small transparent cubic crystals suitable for an X-ray diffraction study were grown from hexane at -30 °C. Complex 5 is monomeric in the solid state. The two cadmium centres are not planar with the ligand, displaced by 0.657 Å for Cd1 and 0.526 Å for Cd2 out of the BDI ligand planes, N1-C1-C2-C3-N2 and N3-C30-C31-C32-N4, respectively. Both cadmium centres are three coordinate in a pseudo-trigonal planar conformation. The N-Cd bond lengths are quite similar and range from 2.197 to 2.212 Å. The N-Cd bonds are larger than those observed in hydride 3 which can be evidence of a change in the oxidation state of the cadmium centres. In 3 the formal oxidation state of the cadmium was Cd(II), in 5 the cadmium centres changed and approach Cd(I) or a [Cd₂]^2+ core, the increase of the ionic radii resulted in the increased bond lengths to the nitrogens of the BDI ligand.

The nitrogen-carbon bonds of the backbone have a small range from 1.310 to 1.330 Å, indicative of a delocalized π-system in the BDI backbones. The two bite angles, N1-Cd1-N2 and N3-Cd2-N4, of 87.14(8)° and 87.46(8)° are similar to hydride 3 and the other three

Figure 15: ORTEP diagram of 5 showing thermal ellipsoids at 30% probability. Isopropyl groups and hydrogens omitted for clarity.
coordinate β-diketiminato-cadmium complexes. The cadmium atoms in 5 have a Cd1-Cd2 bond length of 2.5952(4) Å which is longer than the two coordinate cadmium centres in homonuclear dimer complexes XXI and XXII, which have Cd-Cd bond lengths of 2.574 and 2.579 Å, respectively. The BDI-cadmium centres are not planar to each other and are rotated 74.57° from the other, potentially due to the sterics of the isopropyl groups in the solid state. The ¹H and ¹³C NMR spectra indicate that the ligands at Cd1 and Cd2 are symmetrical in solution as only one set of signals for the BDI ligand is observed.

<table>
<thead>
<tr>
<th>Bond</th>
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<th>Bond</th>
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</table>

The synthesis of complex 5 was only possible upon addition of DCC and DIC to hydride 3, addition of DBC to 3 resulted in a slow degradation of the cadmium hydride to multiple products. The reaction of complex 3 and DDC or DIC (eq 20) did not occur cleanly with minor products being detected throughout the reaction but repeated isolation attempts only yielded complex 5. With smaller equivalents (0.5 and 0.1) of DCC and DIC a cleaner and complete conversion of 3 to 5 was observed. Although small amounts of 5 were observed when 3 degrades over time, other degradation products are not observed during the reaction converting complex 3 to 5. This provides evidence that the reduction of the cadmium hydride is catalytic with DCC or DIC.
Scheme 14: Proposed catalytic production of homonuclear Cd-Cd dimer 5 through amidinate 4 from hydride 3.

This type of reactivity has never been reported with other metal hydride species or carbodiimides. The mechanism for this transformation is unclear, GC-MS (Gas chromatography MS) analysis of the reaction mixture found primarily DCC or DIC in solution, other volatiles were not observed. The initial step could be the insertion of the carbodiimide into the Cd-H bond to give the amidinate 4, this could then decompose to give the homonuclear Cd-Cd dimer (Scheme 14). To do this, we attempted to independently synthesize amidinate 4 via multiple methods.

Cadmium Amidinate Synthesis

Two synthetic methods (A and B) were attempted to synthesize the formamidine starting material (Scheme 15). Method A involved adding a catalytic amount of palladium dichloride and triethylsilane to neat DIC, although this route was reported, we were unable to obtain the desired formamidine. Method B is a known method that involves addition of acetic acid to triethylorthoformate and two equivalents of either neat diisopropylamine or neat cyclohexylamine to form formamidines 6 and 7 respectively. Method B proved successful.

Scheme 15: Methods A and B to generate complexes 6 and 7.
Two methods were attempted to synthesize the cadmium amidinate 4, either via salt metathesis between cadmium chloride complex XVI and lithium amidinate or by treatment of cadmium HMDS complex XV with the formamidine (Scheme 16).

![Scheme 16: Proposed routes to complex 4.](image)

The method using the lithium amidinate was attempted by addition of "BuLi to diisopropyl formamidine 6 in hexane generating the lithium amidinate, which was then added to a toluene solution of chloride XVI. Small white cubic crystals suitable for an X-ray diffraction study from a filtered reaction mixture at -30 °C. These crystals were not the expected amidinate product (4) but were instead a lithium atom coordinating many amidinate molecules in a “caged” structure. The 1H NMR spectrum shows a singlet at δ 8.46 ppm for the proton on the amidinate backbones and a septet at δ 3.38 ppm which indicates a single environment for the central proton on the isopropyl groups of the amidinates.

When cadmium dichloride was used instead of cadmium chloride XVI, a different signal was observed in the 1H NMR spectrum. The 1H NMR spectrum showed multiple products, with a singlet at δ 7.94 ppm with satellites proposed to be 111Cd and 113Cd satellites coupling to the proton of the amidinate backbone. The proposed 113Cd-1H coupling constant, 3J_{HCd}, is 80.5 Hz and 111Cd-1H with 58.5 Hz (Figure 16). The compound was not isolated from the reaction as the lithium “caged” product would preferentially form over time and would crystallize out of the reaction mixture.
The route using cadmium HMDS complex XV was first attempted using [Cd(HMDS)$_2$]. A hexane solution of dicyclohexyl formamidine (7) was added to a solution of [Cd(HMDS)$_2$] synthesized in situ. The reaction mixture yielded a white residue (8).

\[
[Cd(HMDS)_2] + 3 \text{Cy-}N\equiv N\text{-Cy} \xrightarrow{\text{Hex. 48 hr}} \begin{array}{c}
\text{Cy} \\
N \\
\text{Cd} \\
N \\
\text{Cy} \\
\text{Cy} \\
N \\
\text{Cy} \\
\text{Cy} \\
\text{Cy} \\
\text{N} \\
\text{Cy} \\
\text{Cy} \\
\end{array}
\]

(21)

The $^1$H NMR spectrum of 8 shows a singlet at $\delta$ 7.90 ppm, indicating a single environment for the central proton on the amidinate backbone with satellites to cadmium with a coupling constant of 52.4 Hz. The backbone signal is downfield to formamidine 7 ($\delta$ 7.14 ppm). Cyclohexyl peaks are observed as a range of multiplets from $\delta$ 2.04 to 1.28 ppm. Added D$_2$O would not cause degradation and shifted a broad amine hydrogen peak at $\delta$ 3.15 ppm leaving behind a multiplet determined to be the cyclohexyl ipso-carbon’s hydrogen peaks. Transparent rectangular crystals suitable for an X-ray diffraction study were grown from diethyl ether at -30 °C. Although a crystal structure could not fully solved, the connectivity between the cadmium centre and the formamidine ligands were confirmed.
The success with \([\text{Cd(HMDS)}_2]\) gave evidence that the method from cadmium amide complex \(\text{XV}\) would be successful. Attempts to use this method to generate the amidinate complex \(\text{4}\) are ongoing.

Other Reactivity

Addition of phenyl acetylene to hydride \(\text{3}\) resulted in the formation of a black precipitate, \([\text{BDI}_{\text{Dipp}}\text{-H}]\) and \(\text{H}_2\). Addition of isocyanate to hydride \(\text{3}\) resulted in a mixture of products, none of which were isolable. Addition of Super Hydride, potassium hydride or sodium hydride to reduce the cadmium in hydride \(\text{3}\) to generate the homonuclear dimer \(\text{5}\) only formed \([\text{BDI}_{\text{Dipp}}\text{-Li}]\) or \([\text{BDI}_{\text{Dipp}}\text{-K}]\).

Attempts to synthesize \([\text{Ar-BDI}_{\text{Dipp}}\text{-CdCl}]\), a cadmium analogue or mercury chloride \(\text{1}\), were unsuccessful. The salt metathesis route used in the synthesis of complex \(\text{1}\) produced many products with cadmium dichloride, none of which were solid or cleanly isolable. Cadmium halide salts tend to react in ethereal solvents so the route used to synthesis chloride \(\text{XVI}\) was attempted but addition of \([\text{Ar-BDI}_{\text{Dipp}}\text{-H}]\) to \([\text{HMDS-CdCl}]\) did not result in the formation of any isolable products.
2.3 Zinc

2.3.1 Synthesis of a Novel β-Diketiminato-zinc Bromide

As the ligand \([\text{Ar-BDI}_{\text{Dipp}}-\text{H}]\) was able to coordinate to mercury(II), a zinc analogue was synthesized as a comparison. \([\text{Ar-BDI}_{\text{Dipp}}-\text{Li}]\) was generated by adding a solution of \(n\text{BuLi}\) to a toluene solution of \([\text{Ar-BDI}_{\text{Dipp}}-\text{H}]\).\(^{126}\) The intensely yellow precipitate was collected by removing the volatiles \(\text{in vacuo}\); impurities were further removed via triturations in hexane.

The first attempt at generating a β-diketiminato-zinc bromide using the novel BDI involved addition of \([\text{Ar-BDI}_{\text{Dipp}}-\text{Li}]\) to a zinc dibromide slurry in THF. Unfortunately, only a bright yellow oil was isolated, complex 9 (eq 22).

![Chemical structure](image)

The \(^1\text{H} \text{NMR}\) spectrum of 9 shows two septets at \(\delta 3.50\) and \(3.03\) ppm indicating two environments for the methine proton on the isopropyl groups. The proton resonances differ to \([\text{Ar-BDI}_{\text{Dipp}}-\text{Li}]\). Resonances associated with THF are situated at \(\delta 3.57\) and \(1.24\) ppm, consistent with coordinated THF molecules integrating for 16 hydrogens, indicating four THF molecules are incorporated. Although this complex could never be isolated in the solid state, it is suspected that it is similar to the salt metathesis zinc complex XXXII but with more coordinating solvent molecules. This type of reactivity is common with group 12 β-diketiminate complexes.\(^{28, 85}\)

To avoid coordinating solvents, toluene was then used as an alternative. A toluene solution of \([\text{Ar-BDI}_{\text{Dipp}}-\text{Li}]\) was added to a toluene solution of zinc dibromide (eq 23). The reaction retained a strong yellow colour. After 72 hours the reaction mixture was filtered and the volatiles removed \(\text{in vacuo}\) to give a bright yellow solid (10).
The $^1$H NMR spectrum shows two septets at δ 3.34 and 2.83 ppm corresponding to two different environments for the methine proton on the isopropyl groups and four doublets for the isopropyl methyl groups at δ 1.37, 1.31, 1.25 and 0.99, evidence of an asymmetrical ligand environment. The $^1$H NMR resonances differ to [Ar-BDI\textsubscript{dipp}-Li].

Figure 17: ORTEP diagram of 10 bridged dimer showing thermal ellipsoids at 30% probability. Isopropyl groups and hydrogens omitted for clarity.
Small bright yellow cubic crystals suitable for an X-ray diffraction study were grown from hexane at -30 °C. Complex 10 is a bridged dimer in the solid state in which the bromide atoms bridge the zinc centres. The zinc lies out of the BDI backbone plane (N1-C3-C2-C1-N2) by 0.546 Å. The N1-Zn-N2 plane is bent 23.68° from the BDI plane. The backbone aryl group plane consisting of C1-C2-C4-C5-C6-C7 is also bent to a small degree (8.13°) from the BDI plane.

The bridging metallocycle (Zn-Br-Zn’-Br’) is planar, with the Br-Zn-Br’ angle being 89.30(2)° and the Zn-Br-Zn’ angle being 90.70(2)°. The geometry of the zinc metal centre is a distorted tetrahedral with the BDI ligand bite angle of 95.8(1)°. The BDI bite angle is similar to the zinc phosphanide XXX but smaller than reported bridged β-diketiminato-zinc bromide complexes which exhibit a BDI bite angle range of 96.67-98.33° and a bridging bromide bite angle range of 101.26-103.29°.28

The Zn-N1 bond length of 1.998(3) Å is longer than the Zn-N2 bond length of 1.940(2) Å. This is consistent with different types of nitrogen ligands; N1 can be considered an imine, a neutral ligand, whereas N2 can be considered an amide, an anionic ligand. N1 has more imine or double bond character, as observed by the shorter N1-C3 bond length of 1.308(5) Å compared to N2 with a longer N2-C1 bond length of 1.356(5) Å. This asymmetry has been observed with copper complexes bearing this ligand.126 For existing β-diketiminato-zinc complexes bearing asymmetric BDI ligands the amide nitrogens Zn-N bond distance range from 1.907 to 1.949 Å and to the backbone carbon range from 1.340 to 1.374 Å. For the imine nitrogens, the Zn-N bond distance range from 1.978 to 2.044 Å and to the backbone carbon range from 1.289 to 1.318 Å.67-68, 81

The Zn-Br bond is 2.5177(5) Å and the Zn-Br’ bond is 2.4503(5) Å. The zinc-bromide bond lengths are longer than standard zinc dibromide at 2.21 Å124 due to the bridging and electron sharing between two zinc atoms. The Zn-Br bond lengths are similar to reported bridged β-diketiminato-zinc bromides, one bromide bares a shorter bond length ranging 2.402-2.408 Å and one bromide bares longer bond length ranging 2.418-2.435 Å.28
Table 8: Selected bond lengths (Å) and angles (°) for 10.

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<tr>
<th>Bond/Angle</th>
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<th>Bond/Angle</th>
<th>Value</th>
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</thead>
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2.3.2 Derivatization of a Novel β-Diketiminato-zinc Bromide

Synthesis of Amides

Addition of [Li-HMDS] to zinc bromide 10 resulted in the formation of a zinc HMDS complex (11). This complex has a similar intense yellow colour to that of 10 (Scheme 17). The $^1$H NMR spectrum of 11 shows two septets at δ 3.47 and 3.00 ppm corresponding to two different environments for the methine proton on the isopropyl groups and four doublets for the
isopropyl methyl groups at δ 1.45, 1.29, 1.17 and 0.97 ppm. These peaks are evidence of an asymmetrical ligand environment. The $^1$H NMR resonances differ to zinc bromide 10. There is a large singlet at -0.03 ppm integrating for 18 protons for the HMDS substituent, up-field from protonated HMDS and [Li-HMDS].

Bright yellow rectangular plate-like crystals suitable for an X-ray diffraction study were grown from hexane at -30 °C. Complex 11 is a monomer in the solid state and shows two crystallographically identical independent molecules not related by symmetry in the asymmetric unit. The plane consisting of Si1-N6-Si2 is 62.24° to the N1-Zn1-N2 plane potentially due to the crystal packing. Other β-diketiminato-zinc HMDS compounds are all rotated differently to accommodate differing packing and ligand sterics.27, 66, 74, 78, 82

Similar to zinc bromide 10, the two Zn-N bond lengths differ to eachother. The Zn-N1 bond length of 2.017(1) Å is longer than the Zn-N2 bond length of 1.906(1) Å, consistent with different types of nitrogen ligands. N1 can be considered an imine, a neutral ligand, whereas
N2 can be considered an amide, an anionic ligand. N1 has more imine or double bond character due to the shorter N1-C1 bond length of 1.312(2) Å compared to N1 with a longer N2-C3 bond length of 1.362(2) Å.

The Zn-N3 bond length is 1.887(2) Å and the N3-Si bonds are 1.711(2) and 1.715(2) Å. These bond distances are similar to existing β-diketiminato-zinc HMDS compounds which have Zn-N bonds ranging from 1.880 to 1.897 Å and N-Si bond distances ranging from 1.702 to 1.723 Å.²⁷, ⁴⁷, ⁶⁶, ⁷⁴, ⁷⁸, ⁸², ¹³⁸

Table 9: Selected bond lengths (Å) and angles (°) for 11.

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<thead>
<tr>
<th>Bond</th>
<th>Length (Å)</th>
</tr>
</thead>
<tbody>
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<tr>
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<tr>
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<td>1.906(1)</td>
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<td>130.25(6)</td>
</tr>
<tr>
<td>N3-Zn1-N2</td>
<td>134.04(7)</td>
</tr>
</tbody>
</table>

Addition of lithium 2,6-diisopropylphenyl amide (Li-N(H)Dipp) to zinc bromide 10 gives an anilido complex 12. The ¹H NMR spectrum of 12 shows three septets at δ 3.33, 2.93 and 2.54 ppm corresponding to two different environments for the methine proton on the isopropyl groups and one for the NH(Dipp) group, all integrating for two protons. The ¹H NMR spectrum shows four doublets for isopropyl methyl groups at δ 1.26, 1.10, 1.08 and 0.99 ppm which integrate for 6 protons each, denoted as the BDI ligand isopropyl groups, and a doublet at 1.02 ppm integrating for 12 protons, denoted as the NH(Dipp) isopropyl groups. A singlet for the proton off the amide is at 2.96 ppm, integrating for a single proton. This confirms the incorporation of the NH(Dipp) group via a salt metathesis reaction.
Intensely yellow floret-shaped crystals were grown in hexane at room temperature within 12 hours. Despite repeated attempts, the crystals were not suitable for an X-ray diffraction study. This was the first BDI zinc complex to include an aryl amine, other nitrogen containing substituent are either HMDS\textsuperscript{27, 66, 74, 78, 82} or a small amine.\textsuperscript{69, 75}

**Synthesis of Phosphanides**

\[
\text{[Diagram of synthesis reaction]} \quad (24)
\]

\( R = \text{Cy (13)}, \text{Ph (14)} \)

Zinc-phosphanide complexes from lithium phosphanide salts were attempted via a salt metathesis reaction with 11. The lithium phosphanide salts Li-PCy\textsubscript{2} and Li-PPh\textsubscript{2} were used. \( \beta \)-Diketiminato-zinc dicyclohexylphosphanide (13) was synthesized by treatment of zinc bromide 11 with Li-PCy\textsubscript{2} (eq 24). To minimize the possibility of incomplete salt metathesis, a non-ethereal solvent, toluene, was used.

The \(^1\text{H}\) NMR spectrum of 13 shows two septets at \( \delta \) 3.42 and 2.98 ppm corresponding to two different environments for the methine proton on the isopropyl groups and four doublets for the isopropyl methyl groups at \( \delta \) 1.41, 1.38, 1.14 and 0.98 ppm, evidence of an asymmetrical ligand environment. Cyclohexyl peaks are observed in the \(^1\text{H}\) NMR spectrum at \( \delta \) 1.58 ppm and between \( \delta \) 1.32 and 0.98 ppm. The \(^{31}\text{P}\) NMR spectrum shows a single peak at \( \delta \) -22.3 ppm, downfield to Li-PCy\textsubscript{2} with a shift strikingly similar to zinc dicyclohexylphosphanide XXX.
Figure 19: ORTEP diagram of 13 showing thermal ellipsoids at 30% probability. Hydrogens omitted for clarity.

Yellow cube shaped crystals suitable for an X-ray diffraction study were grown from hexane at -30 °C. Complex 13 is monomeric in the solid state and shows disorder at one of the cyclohexyl groups and one of the isopropyl peaks. Both positions were modelled and what are shown in Figure 19 are the nuclear positions with the highest occupancy.

Like the previous compounds bearing the same ligand, there are amide and imine nitrogens of the BDI ligand. The shorter N1-C1 bond (1.312(3) Å) and longer N1-Zn bond (2.018(2) Å) indicates N1 is the imine nitrogen. The longer N2-C3 bond (1.351(4) Å) and shorter N2-Zn bond (1.935(2) Å) indicates N2 is the amide nitrogen. The bite angle (N1-Zn-N2) of 93.13(9)° is similar to the other complexes synthesized bearing the same ligand.

The Zn-P bond length, 2.2822(8) Å, is shorter than a dative phosphanide bond with bond lengths ranging 2.393-2.401 Å\textsuperscript{95} and a P-metallated phosphasilene with a bond length of 2.367 Å.\textsuperscript{97} The Zn1-P1 bond length in complex 13 is closer to that of an internally bound zinc phosphanide in a cluster with bond lengths ranging from 2.302 to 2.334 Å\textsuperscript{96}, like zinc dicyclohexylphosphanide XXX. The phosphanide ligand is asymmetric to the zinc metal.
centre, skewed towards N1 as represented with a shorter N1-Zn-P angle of 127.18(7)° compared to the N2-Zn-P angle of 137.54(7)°. This is observed with zinc dicyclohexylphosphanide XXX to a lesser extent. This is evidence the asymmetric BDI ligand has more of an influence on the terminal phosphanide coordination than a symmetrical BDI ligand.126

Table 10: Selected bond lengths (Å) and angles (°) for 13.

<p>| | | | | | |</p>
<table>
<thead>
<tr>
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Addition of Li-PPh₂ to a toluene solution of zinc bromide 10 produced the zinc diphenylphosphanide complex 14 (eq 24). The ¹H NMR spectrum of 14 shows two septets at δ 3.37 and 2.91 ppm corresponding to two different environments for the methine proton on the isopropyl groups and four doublets for the isopropyl methyl groups at δ 1.13, 1.10, 1.07 and 0.95 ppm, evidence of an asymmetrical ligand environment. Phenyl peaks are observed in the ¹H NMR spectrum as multiple peaks around δ 6.88 ppm. The ³¹P NMR spectrum shows a single peak at δ -47.7 ppm, downfield to Li-PPh₂ and similar to zinc diphenylphosphanide XXXI.
Figure 20: ORTEP diagram of 14 showing thermal ellipsoids at 30% probability. Hydrogens omitted for clarity.

Yellow cube shaped crystals suitable for an X-ray diffraction study were grown from hexane at -30 °C. Complex 14 is a monomer in the solid state and shows two crystallographically identical independent molecules not related by symmetry in the asymmetric unit. Like the previous compounds bearing the same ligand, there are amide and imine nitrogens of the BDI ligand. The shorter N1-C1 bond (1.311(6) Å) and longer N1-Zn1 bond (2.019(4) Å) indicates N1 is the imine nitrogen. The longer N2-C3 bond (1.395(6) Å) and shorter N2-Zn1 bond (1.909(4) Å) indicates N2 is the amide nitrogen. The bite angle (N1-Zn-N2) of 93.6(2)° is similar to dicyclohexylphosphanide 13.

The Zn1-P1 bond length, 2.303(1) Å, is shorter than a dative phosphane bond with bond lengths ranging 2.393-2.401 Å93 and a P-metallated phosphasilene with a bond length of 2.367 Å.97 The Zn1-P1 bond length in complex 14 is closer to that of an internally bound zinc phosphane in a cluster with bond lengths ranging from 2.302 to 2.334 Å95, like the zinc dicyclohexylphosphanide complexes XXX and 13. The phosphane ligand is asymmetric to
the zinc metal centre, skewed towards N1 as represented with a shorter N1-Zn1-P1 angle of 119.4(1)° compared to the N2-Zn1-P1 angle of 146.1(1)°. This is observed with zinc dicyclohexylphosphanide complexes XXX and 13 to a lesser extent. This is further evidence the asymmetric BDI ligand influence on terminal phosphanide coordination.126

**Table 11:** Selected bond lengths (Å) and angles (°) for 14.

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<td>110.2(2)</td>
<td>101.7(2)</td>
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</table>

Addition of Chalcogens into Zinc Phosphanide Bonds

Addition of elemental sulfur to a toluene solution of dicyclohexylphosphanide 13 resulted in the formation of a dicyclohexylphosphinodithioate, complex 15 (Scheme 18). The 1H NMR spectrum of 15 shows two septets at δ 3.65 and 3.15 ppm which correspond to two different environments for the methine proton on the isopropyl groups and four doublets

**Scheme 18:** Synthesis of complexes 15, 16 and 17 from phosphane 13.

Addition of elemental sulfur to a toluene solution of dicyclohexylphosphanide 13 resulted in the formation of a dicyclohexylphosphinodithioate, complex 15 (Scheme 18). The 1H NMR spectrum of 15 shows two septets at δ 3.65 and 3.15 ppm which correspond to two different environments for the methine proton on the isopropyl groups and four doublets
for the isopropyl methyl groups at δ 1.63, 1.59, 1.21 and 0.96 ppm, evidence of an asymmetrical ligand environment. Cyclohexyl peaks are observed in the ¹H NMR spectrum as a range of multiplets from δ 1.55 to 1.13 ppm. The ³¹P NMR spectrum shows a single peak at δ 93.4 ppm, downfield to complex 13. Since the NMR active nuclei of sulfur, ³³S, has a low abundance and sensitivity, satellites in the ³¹P NMR spectrum are not observed. Intensely yellow crystals were grown in hexane at -30 °C.

Despite repeated attempts, the crystals of 15 were not suitable for an X-ray diffraction study. The downfield phosphorus peak in the ³¹P NMR is similar for the other addition complexes XXXIII and XXXIV. The double addition product, a dicyclohexylphosphinodithioate, is the likely product based on the equivalence of elemental sulfur used.

Addition of selenium powder to a toluene solution of dicyclohexylphosphanide 13 resulted in the formation of the dicyclohexylphosphinodiselenoate, complex 16 (Scheme 18). The ¹H NMR spectrum of 16 shows two septets at δ 3.67 and 3.17 ppm corresponding to two different environments for the methine proton on the isopropyl groups and four doublets for the isopropyl methyl groups at δ 1.64, 1.61, 1.23 and 0.98 ppm, evidence of an asymmetrical ligand environment. Cyclohexyl peaks are observed in the ¹H NMR spectrum as a range of multiplets from δ 1.57 to 0.88 ppm. The ³¹P NMR spectrum shows a single peak at δ 51.2 ppm, downfield from dicyclohexylphosphanide 13, with ⁷⁷Se satellites. The ⁷⁷Se-³¹P coupling constant of 447.9 Hz is consistent with the addition of two selenium atoms to a phosphorus atom seen in β-diketiminato-tin and germanium phosphanide complexes.¹²⁵

When the reaction to synthesize complex 16 is monitored by ³¹P NMR spectroscopy, another peak can be observed at δ 25.6 ppm with ⁷⁷Se satellites and a ⁷⁷Se-³¹P coupling constant of 145.8 Hz. The coupling is consistent with the addition of one selenium atom to a phosphorus atom seen in reactions with β-diketiminato-tin and germanium phosphanide complexes.¹²⁵ The signal at δ 25.6 ppm is slowly replaced by the signal of the dicyclohexylphosphinodiselenoate at δ 51.2 ppm over time.
Figure 21: From left to right. 16 with $^{77}$Se satellites, the dicyclohexylphosphinoselenoite product with smaller $^{77}$Se satellites, unreacted lithium dicyclohexylphosphanide and unreacted 13.

It was suspected that the mono-insertion product occurred prior to the addition of a second selenium atom, the equilibrium of which is driven towards the double addition product, dicyclohexylphosphinodiselenoate 16 (Scheme 19). Attempts to isolate the suspected mono-insertion product proved unsuccessful with complex 16 forming preferentially even when treated with a single equivalent or less of selenium powder.

Scheme 19: Selenium insertion reaction with 13 to generate 16.
Orange rectangular crystals suitable for an X-ray diffraction study were grown from hexane at -30 °C. Complex 16 is monomeric in the solid state with a half a molecule of hexane in the unit cell. The phosphorus atom is formally P$^{5+}$ and the selenium atoms are both Se$^{2-}$. The Zn-Se bond lengths of 2.4995(8) and 2.4969(9) Å are very similar where the Se-P bond lengths of 2.166(2) and 2.180(1) Å differ slightly. In contrast to β-diketiminato-germanium dicyclohexylphosphinodiselenoate complexes, which are three coordinate, the dicyclohexylphosphinodiselenoate ligand in complex 16 is bidentate resulting in a tetrahedral zinc centre. The Zn-Se bonds are in range of the Zn-Se bonds in the selenium bridged BDI complex XXIV which range from 2.429-2.595 Å. Other Zn-Se bonds from various compounds range from 2.320-2.609 Å.

Like the previous compounds bearing the same ligand, there are amide and imine nitrogens of the BDI ligand. The shorter N1-C1 bond (1.317(6) Å) and longer N1-Zn bond (2.004(4) Å) indicates N1 is the imine nitrogen. The longer N2-C3 bond (1.350(7) Å) and shorter N2-Zn1 bond (1.947(3) Å) indicates N2 is the amide nitrogen. The bite angle (N1-Zn-N2) of 94.2(2)° is
similar to the other complexes synthesized bearing the same ligand including dicyclohexylphosphanide 13. The plane consisting of N1-C1-C2-C3-N2 is 87.60° from the plane of the phosphinodiselenoate. The phosphinodiselenoate ligand is square shaped, the Zn-Se1-P and Zn-Se2-P bond angles (80.95(4) and 80.74(4)°) are shorter than the Se1-Zn-Se2 and Se1-P-Se2 bond angles (99.77(3) and 108.42(6)°). The bond lengths and angles of the phosphinodiselenoate ligand are similar to zinc phosphinodiselenoate complexes XXXIII and XXXIV.

**Table 12:** Selected bond lengths (Å) and angles (°) for 16.

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<thead>
<tr>
<th>Bond</th>
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</thead>
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<td>C3-N2</td>
<td>1.350(7)</td>
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<tr>
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<tr>
<td>Zn-Se2</td>
<td>2.4969(9)</td>
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<tr>
<td>Se1-P</td>
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<td>1.432(6)</td>
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<td>Zn-N1-C1</td>
<td>124.6(3)</td>
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<td>123.6(5)</td>
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<td>C3-N2-Zn</td>
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<tr>
<td>N2-Zn-Se2</td>
<td>121.1(1)</td>
</tr>
</tbody>
</table>

Addition of tellurium powder to a toluene solution of dicyclohexylphosphanide 13 resulted in the formation of a dicyclohexylphosphinoditellunoate, complex 17 (Scheme 18). The \(^1\)H NMR spectrum of 17 shows two septets at δ 3.74 and 3.27 ppm corresponding to two different environments for the methine proton on the isopropyl groups and three doublets for the isopropyl methyl groups. The isopropyl signal δ 1.64 ppm integrates for 12 protons and 1.29 and 1.05 ppm integrate for 6 protons each, evidence of an asymmetrical ligand environment. Cyclohexyl peaks are observed in the \(^1\)H NMR as a range of multiplets from δ 1.55 to 0.87 ppm. The \(^{31}\)P NMR spectrum shows a single peak at δ 160.3 ppm, downfield to dicyclohexylphosphanide 13. The \(^{125}\)Te-\(^{31}\)P coupling is not observed for this complex and
so the double addition product, a zinc dicyclohexylphosphinoditellunoate, is not confirmed by $^{31}$P NMR spectroscopy.

![ORTEP diagram of 17 showing thermal ellipsoids at 30% probability. Hydrogens and solvent molecule omitted for clarity.](image)

**Figure 23:** ORTEP diagram of 17 showing thermal ellipsoids at 30% probability. Hydrogens and solvent molecule omitted for clarity.

Orange rectangular crystals suitable for an X-ray diffraction study were grown from hexane at -30 °C. Complex 17 is monomeric in the solid state with a half a molecule of hexane in the unit cell. The phosphorus atom is formally P$^{5+}$ and the selenium atoms are both Se$^{2-}$.

The Zn-Te bond lengths of 2.6793(6) and 2.6885(8) Å and the Te-P bond lengths of 2.400(1) and 2.412(1) Å are similar to each other. In complex 17 the Zn-Te bond lengths are shorter than complexes of zinc bearing a telluride which range from 2.573 to 2.581 Å.$^{112, 116, 139}$ The angles between the zinc and the phosphorus (Zn-Te-P) of 80.56(3)$^\circ$ and 80.16(3)$^\circ$ are shorter than reported telluride angles bound to zinc (97.45-103.84$^\circ$)$^{112, 116, 139}$ but are similar to dicyclohexylphosphinodiselenoate 16. In contrast to β-diketiminato-germanium dicyclohexylphosphinoditellunoate complexes, which are three coordinate, the
The BDI ligand has an imine and an amide nitrogen like the previous compounds bearing the same BDI ligand. The shorter N1-C1 bond (1.305(5) Å) and longer N1-Zn bond (2.014(3) Å) indicates N1 is the imine nitrogen. The longer N2-C3 bond (1.353(5) Å) and shorter N2-Zn bond (1.951(3) Å) indicates N2 is the amide nitrogen. The bite angle (N1-Zn-N2) of 93.9(2)° is very similar to dicyclohexylphosphinodiselenoate 16. The plane consisting of N1-C1-C2-C3-N2 is 87.44° from the plane of the phosphinoditellunoate ligand. The phosphinodiselenoate ligand is square shaped, the Zn-Te1-P and Zn-Te2-P bond angles (80.56(3) and 80.16(3)°) are shorter than the Te1-Zn-Te2 and Te1-P-Te2 bond angles (92.01(2) and 106.74(5)°).

**Table 13:** Selected bond lengths (Å) and angles (°) for 17.

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<tr>
<th></th>
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<td>C3-N2</td>
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<tr>
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<td>Zn-Te2</td>
<td>2.6885(8)</td>
<td>Te1-P</td>
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<td>Te2-P</td>
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<td>N2-C21</td>
<td>1.434(5)</td>
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</tr>
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<td>N1-Zn-N2</td>
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<td>Zn-N1-C1</td>
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<td>N1-C1-C2</td>
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<tr>
<td>C1-C2-C3</td>
<td>125.0(4)</td>
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<td>N2-Zn-Te2</td>
<td>121.2(1)</td>
<td></td>
<td></td>
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</table>
Scheme 20: Synthesis of 18 and 19 from 14.

Addition of selenium powder to a toluene solution of diphenylphosphanide 14 resulted in the formation of a diphenylphosphinodiselenoate, complex 18 (Scheme 20). The $^1$H NMR spectrum of 18 shows two septets at $\delta$ 3.78 and 3.27 ppm corresponding to two different environments for the methine proton on the isopropyl groups and four doublets for the isopropyl methyl groups at $\delta$ 1.41, 1.40, 1.26 and 1.03 ppm, evidence of an asymmetrical ligand environment. The $^{31}$P NMR spectrum shows a single peak at $\delta$ 6.1 ppm, downfield from phosphanide 14 with $^{77}$Se satellites.

A $^{77}$Se-$^{31}$P coupling constant of 466.5 Hz is consistent with the addition of two selenium atoms to a phosphorus atom seen in $\beta$-diketiminato-tin and germanium phosphanide complexes. The downfield phosphorus peak and the $^{77}$Se-$^{31}$P coupling in the $^{31}$P NMR are similar to the other phosphinodiselenoate complexes XXXIII, XXXIV and 16. This evidence confirms the diphenylphosphinodiselenoate as the product. Dark yellow crystals were grown in hexane at -30 °C. Despite repeated attempts, the crystals of 18 were not suitable for an X-ray diffraction study.
Addition of tellurium powder to a toluene solution of complex 14 resulted in the formation of multiple products. The $^1$H NMR spectrum of the reaction mixture revealed two septets at $\delta$ 3.40 and 2.85 ppm and four doublets at $\delta$ 1.16, 1.14, 1.07 and 1.03 ppm, evidence of an asymmetrical ligand environment. The $^{31}$P NMR spectrum of the reaction mixture shows a single peak at $\delta$ -15.0 ppm, downfield to Li-PPh$_2$ ($\delta$ -43.9 ppm) and diphenylphosphanide 14. The $^{125}$Te-$^{31}$P coupling is not observed. The crystallized product is a zinc tellunoite bridged dimer (19), which does not contain a phosphorus atom so the signal in the $^{31}$P NMR spectrum is from a by-product.

Phenyl resonances in the $^1$H NMR spectrum are at $\delta$ 7.54-7.48 and 6.94-6.89 ppm as multiplets integrating for 10 protons each. In the $^{13}$C NMR spectrum the peaks not associated with isolated 19 are at $\delta$ 136.61, 134.83, 128.90 and 128.59 ppm. The by-product was proposed to be diphenylidiphosphane, [(Ph$_2$P)$_2$], due to the reported $^{31}$P NMR shift at -15.2 ppm and $^{31}$C NMR shifts of $\delta$ 136.40, 134.86, 128.76 and 129.27.$^{140}$

---

Figure 24: ORTEP diagram of 19 showing thermal ellipsoids at 30% probability. Isopropyl groups, the solvent molecule and hydrogens omitted for clarity.
Golden rectangular plate-like crystals suitable for an X-ray diffraction study were grown from hexane at -30 °C. Complex 19 is monomeric in the solid state with half a molecule of hexane in the unit cell. The tellurium atom is formally Te\(^2-\) with a bent geometry. The Zn1-Te-Zn2 bond angle is 99.09(2)° and has bond lengths to Zn1 and Zn2 of 2.4549(7) and 2.4768(6) Å respectively. In complex 19 the Zn-Te bond lengths are shorter than complexes of zinc bearing a telluride which range from 2.573 to 2.581 Å but the bent angle is within the range of telluride angles involving zinc (97.45-103.84°).\(^{112,116,139}\)

N1 and N3 have longer bonds to their respective zinc centres (1.992(6) and 2.029(4) Å) and shorter bonds to their backbone carbons C1 and C33 (1.307(6) and 1.303(5) Å) making them the imine nitrogens. N2 and N4 have shorter bonds to their respective zinc centres (1.914(4) and 1.910(3) Å) and longer bonds to the backbone carbons C3 and C35 (1.362(6) and 1.374(4) Å) making them the amide nitrogens.

The two BDI ligands are not equivalent and have different bond lengths and angles. The zinc metal centres are displaced from their respective BDI ligand planes (N1-C1-C2-C3-N2 and N3-C33-C34-C35-N4) with Zn1 displaced by 0.426 Å and Zn2 by 0.528 Å. The Zn-Te bonds are skewed, similar to the Zn-P bonds in phosphanide complexes \(^{13,14}\) and XXX. For Zn1, Te is skewed towards N1 as represented with a shorter N1-Zn1-Te angle of 121.5(1)° compared to the N2-Zn1-Te angle of 141.2(1)°. For Zn2, Te is skewed towards N3 as represented with a shorter N3-Zn1-Te angle of 112.6(1)° compared to the N4-Zn1-Te angle of 152.0(1)°. For both centres the Te substituent is skewed towards the imine nitrogen of the BDI ligand, similar to phosphanide complexes \(^{13,14}\) which bare the same ligand. The bite angles differ to each other where the N1-Zn1-N2 angle of 97.1(1)° is larger than previous complexes and the N3-Zn2-N4 angle of 94.2(1)° is similar to the complexes synthesized bearing the same ligand.
**Table 14**: Selected bond lengths (Å) and angles (°) for 19.

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<td>Zn2-N3-C33</td>
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</table>

**Figure 25**: Examples of crystals collected under a microscope. Left, Cubic crystals of dicyclohexylphosphanide 13. Right, Plate-like crystals of telluroite bridged dimer 19.
Absorption occurs when an incoming photon causes an electron to be excited from a lower electronic energy level to a higher electronic energy level. The excited electron then releases energy to return to the original energy level. This energy release can occur through non-radiative processes such as vibrations or through radiative processes which expels a photon, a process called fluorescence. The expelled photon is lower in energy to the photon that caused the absorption, a phenomenon known as the Stokes Shift. Fluorescent molecules often absorb high energy UV light and emit lower energy visible light. An absorption spectrum reveals the wavelengths of photons absorbed over a broad range of light from the UV range through to the visible range.

The photoluminescent properties of molecules can be probed using a spectrofluorophotometer. Two spectra can be obtained from a spectrofluorophotometer, one to determine the wavelengths and intensities of the photons released during fluorescence (an emission spectrum, using a fixed excitation wavelength) and the wavelengths and intensities of the photons being absorbed or causing the excitation of an observed emission (an excitation spectrum, using a fixed emission wavelength). Quantum yield, or fluorescence quantum yield, ($\phi_f$) is a unitless ratio of photons absorbed to photons emitted, providing information on the efficiency of a molecule's fluorescent properties. Emission and excitation intensities and quantum yields have been reported to be influenced by an electron transfer from neighboring lone pairs on heteroatoms to fluorescing $\pi$-systems.141-145

During the characterization of zinc bromide 10, which is intensely yellow in solution, it was found to be fluorescent under a broad range of UV light giving a lime green glow. The amides 11 and 12, the phosphanides 13 and 14, and the addition complexes 15, 16, 17 and 18 were also found to be fluorescent. The $\beta$-diketiminato-zinc complexes XXX, XXXI, XXXIII, and XXXIV bear the BDI$_{Dipp}$ ligand and are not fluorescent; the $\pi$-system of the Ar-BDI$_{Dipp}$ ligand must be the contributing factor.

The fluorescence was occurring due to binding of the BDI ligand to the d$^{10}$ metal ion, zinc,$^{146}$ resulting in an increase in the rigidity of the overall structure compared to the protonated
ligand, [Ar-BDI\textsubscript{Dipp}-H], which is not fluorescent. This gives an extended π-system involving the BDI ligand backbone and the metal centre.\textsuperscript{147} The mercury chloride 1 does not exhibit fluorescent properties potentially due to an insufficient π-system with the mercury metal centre. The orbitals of the metal centre are not directly involved, either from metal-metal transitions or charge transfers from metal to ligand or ligand to metal. This is because it is difficult to oxidize or reduce a highly stabilized full d\textsuperscript{10} configuration, as such found in zinc(II).\textsuperscript{148} The lithium salt, [Ar-BDI\textsubscript{Dipp}-Li], was not fluorescent and did not readily dissolved in solvents used.

Measurements for absorption and photoluminescence were conducted by dissolving a known amount of a sample in hexane, followed by serial dilutions in order to achieve an absorption of approximately 0.1 Abs for the peak around 440 nm. Hexane was chosen as a solvent to avoid solvating processes. The absorption spectra are presented as the extinction coefficient (ε) calculated by Beer-Lambert’s Law (A = c.ε.l).\textsuperscript{149}

An absorption spectrum was collected (Figure 26) followed by an emission spectrum with the fixed excitation wavelength matching the peak found around 440 nm. The emission spectra gave a single broad peak for each compound around 510 nm. The excitation spectra were then collected using the peak around 510 nm for the fixed emission wavelength. The excitation and emission spectra are presented as the intensity normalized by the concentration to allow approximate comparisons (Figure 27).
Quantum yield ($\phi_f$) was calculated for the fluorescent compounds using fluorescein ($\phi_f = 0.18$) in a 2.9 pH citrate-phosphate buffer as a reference, which gives a maximum emission at 510 nm and excitation at 450 nm. The calculation used to determine $\phi_f$ is shown in equation 25. $R$ is the reference compound, fluorescein, $A$ is absorbance at a specified wavelength and $\eta$ is the solvent refractive index.

$$
\phi_f = \phi_f(R) \times \frac{\eta^2}{\eta(R)} \times \frac{1}{L(R)} \times \frac{1 - 10^{-A(R)}}{1 - 10^{-A}} \tag{25}
$$

<table>
<thead>
<tr>
<th>Compound</th>
<th>$\lambda_{max}$ A (nm)</th>
<th>$\varepsilon$ A (M$^{-1}$ cm$^{-1}$)</th>
<th>$\lambda_{max}$ B (nm)</th>
<th>$\varepsilon$ B (M$^{-1}$ cm$^{-1}$)</th>
<th>Excitation (nm)</th>
<th>Emission (nm)</th>
<th>Quantum Yield</th>
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</thead>
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<td>10 (Br)</td>
<td>436</td>
<td>8646</td>
<td>253</td>
<td>25722</td>
<td>434</td>
<td>505</td>
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<tr>
<td>11 (HMDS)</td>
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<td>13389</td>
<td>253</td>
<td>39222</td>
<td>443</td>
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<td>12 (N(H)Dipp)</td>
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<td>10521</td>
<td>253</td>
<td>35014</td>
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These spectra show that the excitation and emissions are not blue or red shifted significantly between each compound, as such, the electronic transitions must be same for each compound. The absorption spectra of the zinc complexes shows two peaks at approximately 440 nm (B) and a higher energy peak at 253 nm (A) (Figure 26) note that the higher energy peak has a shoulder at approximately 290 nm (A*). The literature using this ligand describes similar absorption properties, however the metal centre in the literature, Cu(II), has empty d-orbitals so the transitions were denoted as a π-π* and a d-d transition.\textsuperscript{126} This latter transition cannot occur in the zinc complexes.

The transition around 440 nm in the zinc complexes is proposed to be an n-π* transition from an available lone pair (n) on the substituent off the zinc metal centre (a bromide, amide, phosphanide or chalcogenide) based on reported fluorescent zinc complexes.\textsuperscript{152-154} The n-π* transition occurs via an electron transfer into the π* molecular orbitals of the π-system created when the BDI ligand is coordinated to the zinc centre. The n-π* transition is at a lower energy (B) and the π-π* transition is at a higher energy (A) based on the relative energy between those molecular energy levels.

It is difficult to determine the exact orbitals contributing to fluorescence without a similar system to compare to. This type of system is novel and denotations of the transitions are based on fluorescent zinc molecules that do not contain similar substituents and have different ancillary ligands. The transitions proposed could be confirmed with computational analysis. Density functional theory (DFT) is a computational quantum modelling method used to investigate the electronic structure of systems, including atoms and molecules. Time-dependent density functional theory (TD-DFT) is an extension of DFT. TD-DFT is used to investigate the properties of dynamic systems in the presence of time-dependent processes to extract features including excitation energies and photoluminescence spectra. TD-DFT calculations on the fluorescent zinc complexes could accurately model the transitions involved, the calculations of which are currently ongoing.

The φ\textsubscript{f} are comparable to some zinc complexes involving Schiff bases\textsuperscript{148, 155} but less than zinc salen complexes. This is potentially due to the different coordination and intermolecular aggregations occurring with salen complexes.\textsuperscript{153, 156} Interestingly the ligands used in
reported fluorescent zinc species often exhibit some fluorescence and are enhanced with zinc coordination\textsuperscript{157} In contrast, Ar-BDI\textsubscript{Dipp} is only fluorescent when coordinated to a zinc centre.

One of the least fluorescent compounds is zinc HMDS complex 11. In this complex, the nitrogen lone par has significant back-bonding into the silicon atoms, thus less overlap with the zinc-ligand $\pi$-system would be expected. Although the zinc anilido complex 12 is only mildly fluorescent, anilide ligands typically have less back-bonding of the nitrogen lone pair into the aromatic ring. Quenching of electron transfer has been reported to decrease fluorescence output.\textsuperscript{141-144}

Both of the phosphanide complexes are fluorescent; dicyclohexylphosphanide 13 produces a larger fluorescence intensity compared to diphenylphosphanide 14. The lone pair on the phosphorus in 14 is involved in back-donation into the phenyl substituents, which decreases its $n$-$\pi^*$ transition contribution. The chalcogen addition products, complexes 15 (S), 16 (Se) and 17 (Te), show increasing fluorescence with a decrease in the size of the chalcogen. The two sulfur and selenium atoms have effective orbital overlap to contribute highly in the $n$-$\pi^*$ transition where tellurium has less donation capabilities due to insufficient orbital overlap, resulting in a dramatic loss of fluorescence.

\textbf{Figure 29: Left, samples in ambient light. Right samples under a broad range of UV light. Samples from left to right: Addition of tellurium, selenium and sulfur into the Zn-P bond of complex 13.}
2.4 Summary of Results

Pre-thesis research was conducted using zinc bromide XXIX to synthesize two phosphanides, dicyclohexyl XXX and diphenyl XXXI. The salt metathesis degradation product (XXII) was isolated from the reaction mixture of XXXI. The selenium addition product, dicyclohexylphosphinodiselenoate XXXIII, was synthesized from dicyclohexylphosphanide XXX. The selenium addition product, diphenylphosphinodiselenoate XXXIV, was synthesized from diphenylphosphinide XXXI. The tellurium addition product, dicyclohexylphosphinotellunoite XXXV, was synthesized from dicyclohexylphosphanide XXX and confirmed by 125Te satellites in the 31P NMR spectrum.

The novel β-diketiminato-mercury chloride (1) was synthesized by addition of [Ar-BDI\textsubscript{dipp}-Li], generated by addition of LDA to [Ar-BDI\textsubscript{dipp}-H], to mercury dichloride. Attempts to derivatise 1 using salt metathesis reactions proved unsuccessful with only ligand degradation products being observed.

The recently synthesized β-diketiminato-cadmium chloride (XVI) was derivatized to a β-diketiminato-cadmium phosphanide (2) via a lithium phosphanide, confirmed by Cd\textsuperscript{111} and Cd\textsuperscript{113} satellites in the 31P NMR spectrum, and a novel β-diketiminato-cadmium hydride (3) using Super Hydride. Initial reactivity studies of 3 with various carbodiimides to isolate an insertion product, an amidinate (4), yielded a β-diketiminate with a homonuclear Cd-Cd dimer (5). Complex 5 was formed by catalytic reduction of cadmium hydride 3. Further study is required to confirm the catalytic activity of the carbodiimides and the mechanism with the hydride 3 to produce the homonuclear dimer 5. Attempts to synthesize 4 via a salt metathesis reaction or a ligand exchange reaction from formamidines 6 and 7 proved unsuccessful. A novel cadmium amidinate (8) was synthesized from addition of the formamidine 7 to [Cd(HMDS)\textsubscript{2}].

The same β-diketiminate ligand used to produce mercury chloride 1, [Ar-BDI\textsubscript{dipp}-H], was used to synthesize complexes with zinc. An incomplete salt metathesis reaction product (9) was found when [Ar-BDI\textsubscript{dipp}-Li] was added to zinc dibromide in THF and a novel β-diketiminato-zinc bromide (10) when the reaction was conducted in toluene. Zinc bromide
was derivatized to a variety of complexes (including amides 11 and 12 and phosphanides 13 and 14) by a salt metathesis reaction. Chalcogen addition reactions from dicyclohexylphosphanide 13 gave a zinc dicyclohexylphosphinodithioate (15), dicyclohexylphosphinodiselenoate (16) and a dicyclohexylphosphinoditellunoate (17). Mono-addition products were not isolated. Chalcogen addition reactions from diphenylphosphanide 14 gave a zinc diphenylphosphinodiselenoate (18) and a zinc tellunoite bridged dimer (19). A total of 14 compounds were characterized via X-ray diffraction.

Photoluminescence studies of fluorescent compounds 10-18 were conducted. It was proposed that an electron transfer from the lone pair on the hetero-atom influenced the quantum yield and fluorescence intensities. Further study is required to determine the mechanism of the fluorescence.
Chapter 3

Experimental

All manipulations, unless specified otherwise, were carried out under dry nitrogen using standard Schlenk-line and cannula techniques, or in an inert atmosphere glovebox. Solvents used were dried using a PureSolv. system (Innovative Technologies) and stored over 4 Å sieves unless otherwise stated. $^1$H, $^{13}$C and $^{31}$P NMR spectra were recorded in deuterated benzene (C$_6$D$_6$) at 298 K using a Varian INOVA system at 300 and 600 MHz: ($^1$H), 151 MHz ($^{13}$C ($^1$H)) or 121 MHz ($^{31}$P ($^1$H)). $^1$H and $^{13}$C chemical shifts were referenced internally to residual solvent resonances. Elemental analyses were performed by Campell Micro-analytical Laboratory, Otago. Infrared spectra were recorded on a Bruker Tensor 27 FTIR spectrometer using Nujol mineral oil. Melting points were determined using a Gallenkamp Melting Point Apparatus. Absorption spectra were obtained on a Shimadzu UV-2600 Spectrophotometer. Photoluminescence spectra were obtained on a Shimadzu RF-5301PC Spectrofluorophotometer. Diatomaceous earth used for filtration was Celite™ and is referred to as celite throughout.

Samples suitable for X-ray diffraction were grown in a saturated solution of the sample using at -30 °C unless stated otherwise. Crystals were covered in inert oil and suitable crystals were selected under a microscope and mounted on an Agilent SuperNova X-ray Diffraction system fitted with an Atlas detector. Data were collected at the temperature indicated using focused Mo Kα radiation at 0.71073 Å or Cu Kα radiation at 1.54128 Å. Intensities were collected for Lorentz and polarization effects and for absorption using multi-scan methods. Space groups were determined from systematic absences and checked for higher symmetry. All structures were solved using direct methods with SHELXS, refined on F2 using all data by full matrix least-squares procedures with SHELXL-97, within OLEX-2.3. Non-hydrogen atoms were refined with anisotropic displacement parameters. Hydrogen atoms were placed in calculated positions or manually assigned from residual electron density where appropriate unless otherwise stated. The functions minimized were $\Sigma w(F2o-$
$F2c$, with $w = [\sigma^2(F2o) + aP^2 + bP]^{-1}$, where $P = [\text{max}(Fo)2 + 2F2c]/3$. The isotropic displacement parameters are 1.2 or 1.5 times the isotropic equivalent of their carrier atoms.

The lithium salts of the following: dicyclohexylphosphine, diphenylphosphine, hexamethylsilazane and 2,6-diisopropylphenylamine were synthesized by slow addition of 1.6 M nBuLi in hexane (1.02 mL, 5.04 mmol) to a solution of one of the compounds (5.04 mmol) in hexane (10 mL) for 6 hours. The volatiles were removed in vacuo to yield the product. For lithium dicyclohexylphosphanide, diphenylphosphanide and 2,6-diisopropylphenylamine, triturations in hexane (3 x 5 mL) were conducted to remove impurities. Remaining volatiles were removed in vacuo to reveal the clean lithium salt used throughout.

$[\text{CH}((\text{CH}_3)\text{CN-2,6-Pr}_2\text{C}_6\text{H}_3)_2], \quad [\text{BDI}_\text{Dipp-H}]$ This known compound was synthesized by the following literature procedure. $^{125}$ 12 M HCl (8 mL) was added dropwise to a mixture of acetylacetone (4.7 mL, 45.8 mmol) and freshly distilled 2,6-diisopropylaniline (19.5 mL, 103.4 mmol) in ethanol (200 mL) while stirring. The light pink reaction solution was left to reflux for 72 hours where a milky white precipitate was observed. The precipitate was filtered off and washed with cold petroleum ether (30 mL). Dichloromethane (50 mL) and saturated sodium carbonate solution (100 mL) was added to the solid phase. The aqueous phase was washed further with dichloromethane (50 mL). The organic layers were collected and dried over magnesium sulfate and the volatiles were removed in vacuo. The residue was re-dissolved in methanol (50 mL) and dichloromethane (20 mL) and left to slowly evaporate in a fume cupboard. After 72 hours crystalline product was observed. The product was collected using a Buchner filtration. For use in the inert atmosphere glovebox the crystalline product was crushed and dried at 50 °C under vacuum overnight (14.0 g, 73%). [lit. 73%]. $^{125}$

$^1$H NMR data is in agreement with the literature values. $^1$H NMR (299.74 MHz, $\text{C}_6\text{D}_6$, 298 K): $\delta$ 12.49 (s, 1H, NH), 7.18 (s, 6H, ArH), 4.89 (s, 1H, γ-CH), 3.32 (septet, $J = 6.9$ Hz, $\gamma$, 4H CHMe$_2$), 1.67 (s, 6H, NCMe), 1.23 (d, $J = 7.0$ Hz, 12H, CHMe$_2$), 1.16 (d, $J = 7.0$ Hz, 12H, CHMe$_2$).
[CH{(CH3)CN-2,6-iPr2C6H3}2ZnBr], [BDIDippZnBr] An adapted synthesis was used for this known compound from the following literature procedure.27 [BDIDipp-H] (1.20 g, 2.86 mmol) was dissolved in THF (5 mL) and 1.6 M nBuLi (2.14 mL, 3.43 mmol) was added dropwise to the stirring solution. The reaction was left to react for 30 minutes. Freshly sublimed ZnBr2 (0.77 g, 3.43 mmol) was dissolved in THF (5 mL) and added to the now yellow solution of [BDIDipp-Li] and left to react for 48 hours. The volatiles were removed in vacuo followed by triturations in hexane (3 x 5 mL) to remove impurities. The product was extracted into toluene (12 mL). Toluene and remaining volatiles were removed in vacuo to reveal the product as a white powder. The resulting solid was clean via 1H NMR and used without further purification (1.04 g, 65%). [lit. 96% as chloride derivative].27 1H NMR (299.74 MHz, C6D6, 298 K): δ 7.14 (s, 6H, ArH), 4.89 (s, 1H, γ-C), 3.69 (m, 4H, CHMe2), 1.76 (s, 6H, NCMe), 1.54 (d, J = 3.8 Hz, 12H, CHMe2), 1.22 (d, J = 4.7 Hz, 12H, CHMe2).

[CH{(CH3)CN-2,6-iPr2C6H3}2ZnP(C6H11)2], [BDIDippZnPCy2] (XXX) To a stirring solution of [BDIDippZnBr] (0.200 g, 0.355 mmol) in diethyl ether (5 mL) a lithium dicyclohexylphosphanide (0.073 g, 0.355 mmol) slurry in diethyl ether (10 mL) was added. The reaction was left stirring for 16 hours where a white solid was suspended in a yellow solution. The reaction was filtered through celite and the volatiles were removed in vacuo. The resulting sticky yellow solid was clean via 1H NMR and used without further purification (0.128 g, 52.9 %). Crystals were grown in a minimum amount of diethyl ether. 1H NMR (299.74 MHz, C6D6, 298 K): δ 7.09 (s, 6H, ArH), 5.00 (s, 1H, γ-C), 3.22 (m, 4H, CHMe2), 1.68 (s, 6H, NCMe), 1.56 (m, 18, PCy2), 1.38 (d, J = 6.8 Hz, 12H, CHMe2), 1.28 (m, 6H, PCy2), 1.13 (d, J = 7.1 Hz, 12H, CHMe2). 13C NMR (150.76 MHz, C6D6, 298 K): δ 167.78 (NCMe), 145.79 (ipso-ArC), 141.84 (o-ArC), 126.25 (m-ArC), 124.11 (p-ArC), 95.77 (γ-C), 36.63 (d, JCP = 11.6 Hz, o-PCy), 31.14 (d, JCP = 16.8 Hz, ipso-PCy), 28.70 (CHMe2), 28.45 (d, JCP = 10.4 Hz, m-PCy), 26.41 (CHMe2), 24.36 (d, JCP = 4.6 Hz, p-PCy), 23.93 (CHMe2), 23.78 (NCMe). 31P {1H} NMR (121.32 MHz, C6D6, 298 K): δ -22.2.
[CH(CH$_3$)CN-2,6-$i$Pr$_2$C$_6$H$_3$)$_2$ZnP(C$_6$H$_5$)$_2$, [BDIDippZnPPh$_2$] (XXXI) To a stirring solution of [BDIDippZnBr] (0.260 g, 0.462 mmol) in diethyl ether (5 mL) a lithium diphenylphosphanide (0.089 g, 0.462 mmol) slurry in diethyl ether (10 mL) was added. The reaction was left stirring for 16 hours where a white solid was suspended in a yellow solution. The reaction was filtered through celite and the volatiles were removed in vacuo. The resulting residue was not clean but contained suspected product via $^{31}$P NMR. For use in further reactions the product was made in situ. $^{31}$P $^1$H NMR (121.32 MHz, C$_6$D$_6$, 298 K): $\delta$ -45.8.

[CH(CH$_3$)CN-2,6-$i$Pr$_2$C$_6$H$_3$)$_2$ZnSe$_2$P(C$_6$H$_5$)$_2$, [BDIDippZn(Se)$_2$PCy$_2$] (XXXIII) To a stirring slurry of [BDIDippZnPPh$_2$] (0.120 g, 0.176 mmol) in diethyl ether (5 mL) dried selenium powder was added (0.070 g, 0.882 mmol). The reaction was left to react for 16 hours. The resulting orange solution with dark precipitate was filtered through celite and the volatiles were removed in vacuo. Diethyl ether was added to the residue (2 mL) and kept at -30 °C. Orange rectangular crystals suitable for characterization and X-ray diffraction grew after 72 hours (0.062 g, 42%). $^1$H NMR (299.74 MHz, C$_6$D$_6$, 298 K): $\delta$ 7.22 (s, 6H, ArH), 4.85 (s, 1H, $\gamma$-CH), 3.49 (m, 4H, CHMe$_2$), 1.80 (s, 6H, NCMe), 1.64 (d, J = 7.0 Hz, 12H, CHMe$_2$), 1.50 (m, 16H, PCy$_2$), 1.17 (d, J = 6.7 Hz, 12H, CHMe$_2$), 0.95 (m, 12H, PCy$_2$). $^{13}$C NMR (150.76 MHz, C$_6$D$_6$, 298 K): $\delta$ 167.84 (NCMe), 145.00 (ipso-ArC), 143.34 (o-ArC), 125.62 (m-ArC), 123.91 (p-ArC), 93.44 ($\gamma$-C), 40.05 (d, J$_{CP}$ = 28.3 Hz, ipso-PCy), 28.93 (CHMe$_2$), 26.54 (d, J$_{CP}$ = 2.5 Hz, m-PCy), 26.37 (d, J$_{CP}$ = 14.5 Hz, o-PCy), 26.10 (d, J$_{CP}$ = 2.0 Hz, p-PCy), 25.50 (CHMe$_2$), 25.08 (CHMe$_2$), 24.48 (NCMe). $^{31}$P $^1$H NMR (121.32 MHz, C$_6$D$_6$, 298 K): $\delta$ 51.2 ($J_{PSe}$ = 451.5 Hz).

[CH(CH$_3$)CN-2,6-$i$Pr$_2$C$_6$H$_3$)$_2$ZnSe$_2$P(C$_6$H$_5$)$_2$, [BDIDippZn(Se)$_2$PCy$_2$] (XXXIV) To a stirring slurry of [BDIDippZnPCy$_2$] (0.120 g, 0.176 mmol) in diethyl ether (5 mL) dried selenium powder was added (0.070 g, 0.882 mmol). The reaction was left to react for 16 hours. The resulting orange solution with dark precipitate was filtered through celite and the volatiles were removed in vacuo. Diethyl ether was added to the residue (2 mL) and kept at -30 °C. Orange rectangular crystals suitable for characterization and X-ray diffraction grew after 72 hours (0.062 g, 48%). $^1$H NMR (299.74 MHz, C$_6$D$_6$, 298 K): $\delta$ 7.41 (ddd, J$_{HH}$ = 15.3, 6.9 Hz, J$_{HP}$ = 1.9 Hz, 4H, Ph-m-H), 7.14 (s, 6H, ArH), 6.80 (m, 6H, Ph-o,p-H), 4.84 (s, 1H, $\gamma$-CH), 3.53 (m, 4H, CHMe$_2$), 1.73 (s, 6H, NCMe), 1.41 (d, J = 6.7 Hz, 12H, CHMe$_2$), 1.20 (d, J = 6.7 Hz, 12H, CHMe$_2$). $^{13}$C NMR (150.76 MHz, C$_6$D$_6$, 298 K): $\delta$ 168.30 (NCMe), 144.73 (ipso-ArC), 143.34
\((o\text{-}ArC)\), 136.04 (d, \(J_{CP} = 62.6\) Hz, \(ipso\text{-}PPh\)), 131.06 (d, \(J_{CP} = 12.4\) Hz, \(o\text{-}PPh\)), 130.73 (d, \(J_{CP} = 3.4\) Hz, \(m\text{-}PPh\)), 128.35 (\(p\text{-}PhC\)), 125.93 (\(m\text{-}ArC\)), 124.22 (\(p\text{-}ArC\)), 94.53 (\(\gamma\text{-}C\)), 28.74 (\(\text{C}HMe_2\)), 25.49 (\(\text{CHMe}_2\)), 25.18 (\(\text{CHMe}_2\)), 24.35 (\(\text{NCMe}\)). \(^{31}\text{P} \{^1\text{H}\} \text{NMR (121.32 MHz, C}_6\text{D}_6, 298 K): \delta 6.3 \ (J_{PSe} = 472.7\) Hz).

**[CH\{(\text{CH}_3)\text{CN}-2,6\text{-}i\text{Pr}_2\text{C}_6\text{H}_3\}_2\text{ZnTeP(C}_6\text{H}_{11}\)_2\}, [BDI\text{dipp}Zn(Te)PCy}_2\]** (XXXV) To a stirring slurry of \([\text{BDI}\text{dipp}ZnPCy}_2\] (0.050 g, 0.074 mmol) in diethyl ether (5 mL) dried tellurium powder was added (0.047 g, 0.037 mmol). The reaction was left to react for 48 hours. The resulting dark orange solution with dark precipitate was filtered through celite and the volatiles were removed in \textit{vacuo}. The resulting residue was not clean but contained suspected product via \(^{31}\text{P} \{^1\text{H}\} \text{NMR (121.32 MHz, C}_6\text{D}_6, 298 K): \delta 7.2 \ (J_{PTe} = 285.0\) Hz.

**\([o\text{-}C}_6\text{H}_4\{\text{C(\text{CH}_3)=N-2,6\text{-}i\text{Pr}_2\text{C}_6\text{H}_3\}{\text{F}}\]** An adapted synthesis was used for this known compound from the following literature procedure.\(^{126}\) An ampule fitted with a Young’s tap was charged with TiCl\(_4\) (2.05 g, 0.011 mol) in hexane (15 mL) and 2,6-diisopropylaniline (11.5 g, 0.065 mol) in hexane (15 mL) to form a dense brown precipitate. 2′-fluroacetophenone (2.6 mL, 0.022 mol) was added via syringe which lightened the brown precipitate and allowed to react for 16 hours at 75 °C under N\(_2\). The resulting thick green-yellow solution was opened to air and allowed to cool to room temperature. The contents of the reaction vessel were removed, diethyl ether (3 x 30 mL) was used to remove as much of the solid and liquid from the reaction vessel as possible. The resulting mixture was filtered through celite and a silica plug, the solid fraction was discarded and the orange organic fraction was collected. The orange solution was dried over magnesium sulfate and volatiles were removed in \textit{vacuo} to yield an orange oil. The oil was clean by \(^1\text{H} \text{NMR and used without further purification. This compound was made in situ for the complete synthesis of } [\text{Ar-BDI}\text{dipp-H}] (2.53 \text{ g, 79%})]. \(^{126}\) \(^1\text{H} \text{NMR (299.74 MHz, C}_6\text{D}_6, 298 K): \delta 7.94 \ (td, J = 7.7, 1.9\text{ Hz, } 1\text{H, } \text{ArH}), 7.49 \ (\text{ddd, J = 8.1, 7.3, 5.3 Hz, } 1\text{H, } \text{ArH}), 7.31-7.08 \ (m, 5\text{H, } \text{ArH}), 2.86 \ (\text{septet, J = 6.9 Hz, } 2\text{H, CHMe}_2), 2.17 \ (d, J_{HF} = 3.5\text{ Hz, } 3\text{H, NCMe}), 1.32 \ (d, J = 6.9\text{ Hz, } 6\text{H, CHMe}_2), 1.22 \ (d, J = 6.8\text{ Hz, } 6\text{H, CHMe}_2).
[\text{o-C}_6\text{H}_4{\text{C(CH}_3)=\text{N}-2,6-}\text{iPr}_2\text{C}_6\text{H}_3]\{\text{NH}(2,6-\text{iPr}_2\text{C}_6\text{H}_3)\}], \text{[Ar-BDIDipp-H]} \text{An adapted synthesis was used for this known compound from the following literature procedure.}^{126} 1.6 \text{ M } \text{nBuLi (7.4 mL, 0.012 mol) in hexane was added to 2,6-diisopropylaniline (2.30 g, 0.013 mmol) in THF (12 mL). The reaction was allowed to react for 3 hours.} \text{[o-C}_6\text{H}_4{\text{C(CH}_3)=\text{N}-2,6-}\text{iPr}_2\text{C}_6\text{H}_3]\{\text{F}\} \text{was made in situ (above), was dissolved in THF (12 mL) and added to the lithium 2,6-diisopropylanilide reaction. The total reaction was allowed to react at 90 °C for 72 hours under N}_2. \text{The dark and yellow solution was allowed to cool to room temperature and quenched with H}_2\text{O (30 mL) and extracted into hexanes (50 mL). The aqueous fraction was further washed with hexanes (2 x 20 mL). The organic fractions were combined and dried over magnesium sulfate. Volatiles were removed in vacuo to yield an oily orange solid. The residue was re-dissolved in methanol (30 mL) and dichloromethane (10 mL) and left to slowly crystallize in a fume hood. Yellow crystalline product was observed within 72 hours (3.66 g, 67%). [lit. 46%].}^{126} \text{1H NMR (299.74 MHz, C}_6\text{D}_6, 298 K): } \delta 11.58 \text{ (s, 1H, NH)}, 7.48 \text{ (d, J = 8.2 Hz, 1H, C}_6\text{H}_4), 7.28-7.18 \text{ (m, 6H, ArH), 7.00} \text{ (t, J = 7.6 Hz, 1H, C}_6\text{H}_4), 6.58 \text{ (t, J = 7.5 Hz, 1H, C}_6\text{H}_4), 6.48 \text{ (d, J = 8.5 Hz, 1H, C}_6\text{H}_4), 3.44 \text{ (septet, J = 7.3 Hz, 2H, CHMe}_2), 3.05 \text{ (septet, J = 7.0 Hz, 2H, CHMe}_2), 1.96 \text{ (s, 6H, NCMMe), 1.17} \text{ (d, J = 2.1 Hz, 6H, CHMe}_2), 1.13 \text{ (d, J = 7.0 Hz, 6H, CHMe}_2).}

[\text{o-C}_6\text{H}_4{\text{C(CH}_3)=\text{N}-2,6-}\text{iPr}_2\text{C}_6\text{H}_3]\{\text{NLi}(2,6-\text{iPr}_2\text{C}_6\text{H}_3)\}], \text{[Ar-BDIDipp-Li]} \text{An adapted synthesis was used for this known compound from the following literature procedure.}^{126} 1.6 \text{ M } \text{nBuLi in hexanes (0.62 mL, 0.992 mmol) was added slowly to a stirring solution of} \text{[Ar-BDIDipp-H]} \text{(0.410 g, 0.902 mmol) in toluene (10 mL). The solution immediately turned orange from yellow. A bright yellow precipitate was observed within 4 hours. The reaction was left to react for 16 hours. The volatiles were removed in vacuo followed by triturations in hexane (2 x 10 mL) to further remove impurities. The bright yellow solid was clean by } ^1\text{H NMR and was used without further purification (0.35 g, 84%). [lit. 86%].}^{126} \text{1H NMR (299.74 MHz, C}_6\text{D}_6, 298 K): } \delta 7.63 \text{ (d, J = 8.8 Hz, 1H, C}_6\text{H}_4), 7.32-7.10 \text{ (m, 6H, ArH), 7.03} \text{ (t, J = 7.3 Hz, 1H, C}_6\text{H}_4), 6.55 \text{ (d, 1H, J = 9.0 Hz, C}_6\text{H}_4), 6.35 \text{ (t, 1H, J = 7.1 Hz, C}_6\text{H}_4), 3.31 \text{ (m, 2H, CHMe}_2), 2.78 \text{ (m, 2H, CHMe}_2), 2.06 \text{ (s, 6H, NCMMe), 1.21} \text{ (d, J = 6.7 Hz, 3H, CHMe}_2), 1.17 \text{ (d, J = 6.7 Hz, 3H, CHMe}_2), 1.06 \text{ (d, J = 6.7 Hz, 3H, CHMe}_2), 1.03 \text{ (d, J = 6.7 Hz, 3H, CHMe}_2). \text{The reaction was repeated using lithium diisopropylamide (0.106 g, 0.992 mmol) dried from a slurry in hexane to yield a}
slightly darker yellow solid. This sample had shifted peaks and was used for use with HgCl₂.
(0.34 g, 83%). 1H NMR (299.74 MHz, C₆D₆, 298 K): δ 7.58 (d, J = 7.0 Hz, 1H, C₆H₄), 7.32-7.04
(m, 6H, ArH), 6.58 (d, J = 6.9 Hz, 1H, C₆H₄), 6.35 (t, J = 7.0 Hz, 1H, C₆H₄), 3.54 (septet, J = 6.8 Hz, 2H, CHMe₂), 3.10 (septet, J = 7.1 Hz, 2H, CHMe₂), 2.01 (s, 6H, NCMe), 1.32 (d, J = 7.0 Hz, 3H, CHMe₂), 1.16 (d, J = 7.1 Hz, 3H, CHMe₂), 1.14 (d, J = 7.0 Hz, 3H, CHMe₂), 1.11 (d, J = 7.0 Hz, 3H, CHMe₂).

[α-C₆H₄(C(CH₃)=N{-2,6-Pr₂C₆H₃})₂HgCl], [Ar-BDIDippHgCl] (1) To a stirring solution of [Ar-BDIDipp-Li] (0.140 g, 0.304 mmol) in toluene (10 mL) at -30 °C, HgCl₂ (0.083 g, 0.304 mmol) was added dry. The reaction was left to react for 24 hours where a white solid was suspended in a light yellow solution. The reaction was filtered through celite and the volatiles were removed in vacuo. The resulting light yellow solid was clean via 1H NMR (0.040 g, 69%). Light yellow flower shaped crystals was grown in hexane. 1H NMR (299.74 MHz, C₆D₆, 298 K): δ 7.29 (d, 1H, J = 8.0 Hz, C₆H₄), 6.54 (d, 1H, J = 8.7 Hz, C₆H₄), 3.48 (septet, 2H, J = 6.9 Hz, CMe₂), 1.89 (s, 6H, NCMe), 1.33 (d, 6H, J = 6.9 Hz, CHMe₂), 1.24 (d, 6H, J = 7.0 Hz, CHMe₂), 1.14 (d, 6H, J = 6.9 Hz, CHMe₂), 1.04 (d, 6H, J = 6.8 Hz, CHMe₂). 13C NMR (150.76 MHz, C₆D₆, 298 K): δ 175.04 (NCMe), 155.86 (ArC), 145.93 (ArC), 145.10 (ArC), 137.94 (ArC), 132.69 (ArC), 131.80 (ArC), 126.83 (ArC), 126.19 (ArC), 125.19 (ArC), 124.18 (ArC), 122.18 (ArC), 120.48 (ArC), 116.56 (ArC), 28.86 (CHMe₂), 27.83 (CHMe₂), 24.78 (CHMe₂), 24.72 (CHMe₂), 24.52 (CHMe₂), 24.00 (CHMe₂), 23.34 (NCMe).

(CH{(CH₃)CN{-2,6-Pr₂C₆H₃}}₂CdCl], [BDIDippCdCl] (XVI) An adapted synthesis was used for this known compound from the following literature procedure. Lithium hexamethyldisilazane (0.830 g, 4.96 mmol) was dissolved in 5 mL of THF and added drop-wise to a slurry of CdCl₂ (0.910 g, 4.96 mmol) in THF (5 mL). The reaction mixture was left to stir 16 hours where a grey solution had appeared. The volatiles were removed in vacuo to yield a thick grey oil. The oil was re-dissolved in toluene and filtered through celite. This solution contained (HMDS)CdCl. A solution of BDIDippH (1.20 g, 2.86 mmol) in toluene was added to the (HMDS)CdCl solution and left to react for 72 hours at 55 °C. The volatiles were removed in vacuo followed by triturations in hexane (3 x 10 mL) to remove impurities. Hexane was
evaporated under reduced pressure followed by [BDIDi²pCdCl] extraction into toluene (3 x 10 mL). Removing volatiles in vacuo gave the product as a white powder (0.283 g, 18%). [lit. 57%].

$^1$H NMR (299.74 MHz, $\text{C}_6\text{D}_6$, 298 K): δ 7.13 (s, 6H, ArH), 4.64 (s, 1H, γ-CH), 3.23 (septet, $J = 6.9$ Hz, 4H, CHMe$_2$), 1.56 (s, 6H, NCMe), 1.14 (d, $J = 7.0$ Hz, 12H, CHMe$_2$), 1.01 (d, $J = 6.7$ Hz, 12H, CHMe$_2$).

**[CH{(CH$_3$)CN-2,6-iPr$_2$C$_6$H$_3$}$_2$CdP(C$_6$H$_{11}$)$_2$], [BDIDi²pCdPCy$_2$]** (2) To a stirring solution of [BDIDi²pCdCl] (0.200 g, 0.350 mmol) in toluene (5 mL) a lithium dicyclohexylphosph anide (0.072 g, 0.350 mmol) slurry in toluene (10 mL) was added. The reaction was left stirring for 16 hours where a white solid was suspended in a yellow solution. The reaction was filtered through celite and the volatiles were removed in vacuo. The resulting yellow solid was washed with hexane. A product was not cleanly isolated but suspected product peaks were observed via $^1$H and $^{31}$P NMR. $^1$H NMR (299.74 MHz, $\text{C}_6\text{D}_6$, 298 K): δ 7.06 (s, 6H, ArH), 4.92 (s, 1H, γ-CH), 3.31 (m, 4H, CHMe$_2$), 1.72 (s, 6H, NCMe), 1.35 (d, $J = 6.8$ Hz, 12H, CHMe$_2$), 1.16 (d, $J = 6.7$ Hz, 12H, CHMe$_2$), cyclohexyl peaks were a range of multiplets from 1.86-0.78 ppm. $^{31}$P ($^1$H) NMR (121.32 MHz, $\text{C}_6\text{D}_6$, 298 K): δ 5.4 (J$_{p^{\text{m}}\text{Cd}} = 545.8$ Hz, J$_{p^{\text{n}}\text{Cd}} = 570.8$ Hz)

**[CH{(CH$_3$)CN-2,6-iPr$_2$C$_6$H$_3$}$_2$CdH], [BDIDi²pCdH]** (3) 238 μL of LiHBEt$_3$ in THF (0.0252 g, 0.238 mmol) was added drop-wise to a slurry of [BDIDi²pCdCl] (0.150 g, 0.265 mmol) in 4 mL of toluene at -30 °C. The mixture was allowed to stir for 1 minute where it was filtered through celite to remove black precipitate and the volatiles were removed in vacuo. Solid was re-dissolved and crystallized from hexane or diethyl ether to give [BDIDi²pCdH] as small colourless cubic crystals (0.0944 g, 74.6%). The product was stable at room temperature but decomposed above 126 °C. $^1$H NMR (299.74 MHz, $\text{C}_6\text{D}_6$, 298 K): δ 7.11 (s, 6H, ArH), 5.63 (br, 1H, CdH), 4.91 (s, 1H, γ-CH), 3.29 (septet, $J = 6.7$ Hz, 4H, CHMe$_2$), 1.70 (s, 6H, NCMe), 1.23 (d, $J = 6.8$ Hz, 12H, CHMe$_2$), 1.16 (d, $J = 6.7$ Hz, 12H, CHMe$_2$). $^{31}$C ($^1$H) NMR (150.76 MHz, $\text{C}_6\text{D}_6$, 298 K): δ 168.14 (NCMe), 146.42 (ipso-Ç), 141.10 (o-Ç), 125.76 (p-Ç), 124.00 (m-Ç), 94.71 (γ-Ç), 28.21 (CHMe$_2$), 24.79 (CHMe$_2$), 24.10 (CHMe$_2$), 23.30 (NCMe). IR (Nujol): ν(Cd-H) 1734 cm$^{-1}$
[CH{(CH\textsubscript{3})CN-2,6-\textit{i}Pr\textsubscript{2}C\textsubscript{6}H\textsubscript{3}}\textsubscript{2}Cd]\textsubscript{2}, \ [\text{BDIDippCd}]\textsubscript{2} (5) To a solution of [\text{BDIDippCdH}] (0.050 g, 0.094 mmol) in hexane (5 mL) a solution of dicyclohexylcarbodiimide (0.019 g, 0.094 mmol) in hexane (5 mL) was added. The colourless reaction was left to react for 16 hours. Volatiles were removed in \textit{vacuo} from the colourless and transparent solution. The resulting residue was re-dissolved into hexane (2 mL) and kept at -30 °C. Transparent colourless cubic crystals suitable for X-ray diffraction were grown within 24 hours (0.078 g, 78%). The same procedure was used with diisopropylcarbodiimide and smaller equivalents of either yielded the same results. \textit{\textsuperscript{1}H} NMR (299.74 MHz, C\textsubscript{6}D\textsubscript{6}, 298 K): δ 7.04 (s, 6H, ArH), 4.78 (s, 1H, γ-CH), 3.12 (septet, J = 6.9 Hz, 4H, CHMe\textsubscript{2}), 1.57 (s, 6H, NCMe), 1.16 (d, J = 6.7 Hz, 12H, CHMe\textsubscript{2}), 0.92 (d, J = 7.0 Hz, 12H, CHMe\textsubscript{2}). \textit{\textsuperscript{13}C} NMR (150.76 MHz, C\textsubscript{6}D\textsubscript{6}, 298 K): δ 166.74 (NCMe), 147.79 (ipso-C), 141.06 (ω-C), 125.02 (β-C), 123.88 (m-C), 95.04 (γ-C), 28.24 (CHMe\textsubscript{2}), 24.94 (CHMe\textsubscript{2}), 24.63 (CHMe\textsubscript{2}), 23.06 (NCMe).

\[[\text{CH(N-C\textsubscript{6}H\textsubscript{11})\textsubscript{2}}\textsubscript{2}\text{CH(N-C\textsubscript{6}H\textsubscript{11})(N(H)-C\textsubscript{6}H\textsubscript{11})}]\text{Cd}\], \ [[\text{DC-Am})\textsubscript{2}(\text{DC-Fo})\text{Cd}] \quad (8) \] Lithium hexamethyldisilazane (0.420 g, 2.51 mmol) was dissolved in 5 mL of THF and added drop-wise to a slurry of CdCl\textsubscript{2} (0.230 g, 1.23 mmol) in THF (5 mL). The reaction mixture was left to stir 16 hours where an orange solution had appeared. The volatiles were removed in \textit{vacuo} to yield a thick orange-grey oil. The oil was re-dissolved in toluene (5 mL) and filtered through celite. A solution of DC-Fo (0.52 g, 2.51 mmol) in hexane was added to the Cd(HMDS)\textsubscript{2} solution and left to react for 48 hours. The volatiles were removed in \textit{vacuo} yielding a thick yellow oil. Diethyl ether was added to the residue (2 mL) and kept at -30 °C. Transparent rectangular crystals suitable for characterization and X-ray diffraction grew after 24 hours (0.47 g, 52%) \textit{\textsuperscript{3}H} NMR (299.74 MHz, C\textsubscript{6}D\textsubscript{6}, 298 K): δ 7.90 (s, 3J\textsubscript{H-Cd} = 52.4 Hz, 3H, NC(H)N), 3.15 (br, NC(H)N-H), 2.01 (m, NCy), 1.85 (m, NCy), 1.59 (m, NCy), 1.31 (m, NCy).

\[[\text{Ar}\text{ BDIDippZnBr}_2\text{Li(THF)}_4]\] \quad (9) To a stirring solution of ZnBr\textsubscript{2} (0.034 g, 0.152 mmol) in THF (5 mL), [\text{Ar-BDI\textsubscript{dipp}ZnBr\textsubscript{2}Li(THF)}\textsubscript{4}] (0.070 g, 0.152 mmol) in THF (5 mL) was added dropwise. The reaction was left to stir for 72 hours where a yellow precipitate was suspended in a yellow solution. The reaction was filtered through celite and the volatiles were removed in \textit{vacuo}. The bright yellow sticky oil was clean via \textit{\textsuperscript{1}H} NMR (0.132 g, 89%). Further removal of solvent proved impossible. \textit{\textsuperscript{1}H} NMR
(299.74 MHz, C₆D₆, 298 K): δ 7.30 (dd, 1H, J = 8.5, 1.5 Hz, C₆H₄), 7.21-6.99 (s, 6H, ArH), 6.78 (td, J = 6.9, 1.8 Hz, 1H, C₆H₄), 6.53 (dd, J = 8.7, 1.2 Hz, 1H, C₆H₄), 6.30 (td, J = 6.5, 1.5 Hz, 1H, C₆H₄), 3.57 (m, 16H, C₆H₄Me₂), 3.50 (septet, J = 6.7 Hz, 2H, C₆H₄Me₂), 1.94 (s, 6H, ArH), 1.39 (d, J = 6.8 Hz, 3H, CH₃Me₂), 1.37 (d, J = 6.7 Hz, 3H, CH₃Me₂), 1.24 (m, 16H, CH₃Me₂), 0.95 (d, J = 6.8 Hz, 6H, CH₃Me₂).

[0-C₆H₄(C(CH₃)=N-2,6-iPr₂C₆H₃){NH(2,6-iPr₂C₆H₃)}ZnBr], [Ar-BDI_{dipp}ZnBr] (10) To a stirring slurry of sublimed ZnBr₂ (0.068 g, 0.304 mmol) in toluene (10 mL), [Ar-BDI_{dipp}-Li] (0.140 g, 0.304 mmol) in toluene (5 mL) was added dropwise. The reaction was left to stir for 72 hours where a yellow precipitate was suspended in a yellow solution. The reaction was filtered through celite and the volatiles were removed in vacuo. The bright yellow product was clean via ¹H NMR (0.150 g, 82%). Bright yellow cubic crystals were grown in minimum hexane. ¹H NMR (299.74 MHz, C₆D₆, 298 K): δ 7.44 (d, J = 6.6 Hz, 1H, C₆H₄), 7.24 (s, 3H, ArH), 7.08 (m, 3H, ArH), 6.88 (t, J = 7.3 Hz, 1H, C₆H₄), 6.69 (d, J = 9.1 Hz, 1H, C₆H₄), 6.40 (t, J = 7.8 Hz, 1H, C₆H₄), 3.34 (septet, J = 6.8 Hz, 2H, CH₃Me₂), 2.83 (septet, J = 6.9 Hz, 2H, CH₃Me₂), 1.99 (s, 6H, NC₆Me), 1.37 (d, J = 6.8 Hz, 3H, CH₃Me₂), 1.31 (d, J = 6.7 Hz, 3H, CH₃Me₂), 1.13 (d, J = 7.0 Hz, 3H, CH₃Me₂), 0.95 (d, J = 6.8 Hz, 6H, CH₃Me₂). ¹³C NMR (150.76 MHz, C₆D₆, 298 K): δ 177.86 (NC₆Me), 159.15 (ArC), 144.00 (ArC), 143.07 (ArC), 142.38 (ArC), 139.95 (ArC), 134.36 (ArC), 133.89 (ArC), 127.62 (ArC), 126.57 (ArC), 124.74 (ArC), 124.53 (ArC), 118.66 (ArC), 114.82 (ArC), 114.44 (ArC), 29.18 (CH₃Me₂), 28.59 (CH₃Me₂), 24.92 (CH₃Me₂), 24.28 (CH₃Me₂), 24.18 (CH₃Me₂), 23.64 (CH₃Me₂), 21.51 (NC₆Me). UV-vis (Hexane, 25 °C) [λ_{max}, nm (ε, mol⁻¹ L cm⁻¹)]: 253 (25,700), 435 (8,600).

[0-C₆H₄(C(CH₃)=N-2,6-iPr₂C₆H₃){NH(2,6-iPr₂C₆H₃)}ZnN(SiMe₃)₂], [Ar-BDI_{dipp}ZnHMDS] (11) To a stirring solution of [Ar-BDI_{dipp}ZnBr] (0.050 g, 0.084 mmol) in hexane (5 mL) a lithium Hexamethyldisilazane (0.014 g, 0.084 mmol) solution in hexane (5 mL) was added. The reaction was left stirring for 16 hours where a white solid was suspended in a yellow solution. The reaction was filtered through celite and the volatiles were removed in vacuo. The resulting yellow solid was clean via ¹H NMR (0.040 g, 70%). Rectangular plate-like crystals were grown in hexane. ¹H NMR (299.74 MHz, C₆D₆, 298 K): δ 7.33 – 7.25 (m, 4H), 7.11 (t, J = 1.9 Hz, 3H), 6.83 (ddd, J = 8.6, 6.6, 1.6 Hz, 1H, C₆H₄), 6.60 (dd, J = 8.8, 1.2 Hz, 1H,
C₆H₄), 6.35 (ddd, J = 8.2, 6.7, 1.2 Hz, 1H, C₆H₄), 3.47 (septet, J = 6.8 Hz, 2H, CHMe₂), 3.00 (septet, J = 6.7 Hz, 2H, CHMe₂), 1.93 (s, 3H, NCMe), 1.45 (d, J = 6.9 Hz, 6H, CHMe₂), 1.29 (d, J = 6.7 Hz, 6H, CHMe₂), 1.17 (d, J = 6.8 Hz, 6H, CHMe₂), 0.97 (d, J = 6.7 Hz, 6H, CHMe₂), -0.03 (s, 18H, N(SiMe₃)₂). ¹³C NMR (150.76 MHz, C₆D₆, 298 K): δ 178.88 (NCMe), 158.42 (ArC), 145.54 (ArC), 144.62 (ArC), 144.54 (ArC), 140.18 (ArC), 132.90 (ArC), 127.34 (ArC), 125.97 (ArC), 124.95 (ArC), 124.92 (ArC), 120.73 (ArC), 116.91 (ArC), 113.78 (ArC), 113.78 (ArC), 127.40 (ArC), 126.45 (ArC), 125.24 (ArC), 124.71 (ArC), 122.54 (ArC), 119.24 (ArC), 116.57 (ArC), 115.71 (ArC), 114.15 (ArC), 29.28 (CHMe₂), 28.76 (CHMe₂), 28.51 (CHMe₂), 24.61 (CHMe₂), 23.89 (CHMe₂), 23.77 (CHMe₂), 23.71 (CHMe₂), 23.46 (CHMe₂), 22.06 (NCMe). UV-vis (Hexane, 25 °C) [λmax, nm (ε, mol⁻¹ L cm⁻¹)]: 253 (35,000), 437 (10,500).

[ο-C₆H₄{C(CH₃)=N-2,6-iPr₂C₆H₃}{NH(2,6-iPr₂C₆H₃)}ZnNH(2,6-iPr₂C₆H₃)], [Ar-BDiPpZnNH(Dipp)] (12) To a stirring solution of [Ar-BDiPpZnBr] (0.050 g, 0.084 mmol) in hexane (5 mL) a lithium 2,6-diisopropylaniline (0.015 g, 0.084 mmol) solution in hexane (5 mL) was added. The reaction was left stirring for 16 hours where a white solid was suspended in a yellow solution. The reaction was filtered through celite and the volatiles were removed in vacuo. The resulting yellow solid was clean via ¹H NMR (0.046 g, 82%). A bright yellow flower shaped crystals were grown in hexane but proved inappropriate for X-ray diffraction. ¹H NMR (299.74 MHz, C₆D₆, 298 K): δ 7.41 (dd, J = 8.5, 1.6 Hz, 1H, C₆H₄), 7.21 (s, 3H, ArH), 7.09 (m, 3H, ArH), 7.04 (d, J = 7.6 Hz, 2H, DippArH), 6.83 (dd, J = 8.4, 7.0 Hz, 2H, C₆H₄), 6.80 (d, J = 7.4 Hz, 1H, DippArH), 6.55 (dd, J = 9.0, 1.2 Hz, 1H, C₆H₄), 6.39 (ddd, J = 8.1, 6.6, 1.2 Hz, 1H, C₆H₄), 3.33 (septet, J = 6.9 Hz, 2H, CHMe₂), 2.96 (s, 1H, DippNH), 2.93 (septet, J = 6.9 Hz, 2H, CHMe₂), 2.54 (septet, J = 6.7 Hz, 2H, CHMe₂), 1.93 (s, 3H, NCMe), 1.26 (d, J = 6.9 Hz, 6H, CHMe₂), 1.10 (d, J = 7.0 Hz, 6H, CHMe₂), 1.08 (d, J = 6.9 Hz, 6H, CHMe₂), 1.02 (d, J = 6.7 Hz, 12H, DippCHMe₂), 0.99 (d, J = 6.8 Hz, 6H, CHMe₂). ¹³C NMR (150.76 MHz, C₆D₆, 298 K): δ 177.03 (NCMe), 158.70 (ArC), 148.85 (ArC), 145.08 (ArC), 144.09 (ArC), 143.38 (ArC), 140.04 (ArC), 134.91 (ArC), 133.72 (ArC), 133.37 (ArC), 127.40 (ArC), 126.45 (ArC), 125.24 (ArC), 124.71 (ArC), 122.54 (ArC), 119.24 (ArC), 116.57 (ArC), 115.71 (ArC), 114.15 (ArC), 29.28 (CHMe₂), 28.76 (CHMe₂), 28.51 (CHMe₂), 24.61 (CHMe₂), 23.89 (CHMe₂), 23.77 (CHMe₂), 23.71 (CHMe₂), 23.46 (CHMe₂), 22.06 (NCMe). UV-vis (Hexane, 25 °C) [λmax, nm (ε, mol⁻¹ L cm⁻¹)]: 253 (39,200), 443 (13,400).
[o-C₆H₄{C(CH₃)=N-2,6-iPr₂C₆H₃}{NH(2,6-iPr₂C₆H₃)}ZnP(C₆H₁₁)₂], [Ar-BDIDippZnPPh₂] (13) To a stirring solution of [Ar-BDIDippZnBr] (0.80 g, 0.134 mmol) in hexane (5 mL) a lithium dicyclohexylphosphanide (0.027 g, 0.134 mmol) slurry in hexane (5 mL) was added. The reaction was left stirring for 36 hours where a white solid was suspended in a yellow solution. The reaction was filtered through celite and the volatiles were removed in vacuo. The resulting yellow solid was clean via ¹H NMR and used without further purification (0.059 g, 61%). Yellow cubic crystals were grown from hexane. ¹H NMR (299.74 MHz, C₆D₆, 298 K): δ 7.49 (d, J = 8.5 Hz, 1H, C₆H₄), 7.29-7.02 (m, 6H, Ar H), 6.89 (t, J = 7.8 Hz, 1H, C₆H₄), 6.64 (d, J = 8.9 Hz, 1H, C₆H₄), 6.40 (t, J = 7.7 Hz, 1H, C₆H₄), 3.42 (septet, J = 6.8 Hz, 2H, C₇HMe₂), 2.98 (septet, J = 6.8 Hz, 2H, C₇HMe₂), 2.00 (s, 6H, NCMe), 1.58 (m, 8H, PCy₂), 1.41 (d, J = 6.9 Hz, 6H, CHMe₂), 1.38 (d, J = 6.8 Hz, 6H, CHMe₂), 1.14 (d, J = 6.9 Hz, 6H, CHMe₂), 0.98 (d, J = 6.8 Hz, 6H, CHMe₂), other cyclohexyl peaks were a range of multiplets from 1.32-0.98 ppm.

¹³C NMR (150.76 MHz, C₆D₆, 298 K): δ 175.15 (NCMe), 158.03 (Ar C), 146.13 (Ar C), 145.48 (Ar C), 144.24 (Ar C), 140.03 (Ar C), 133.45 (Ar C), 133.40 (Ar C), 127.09 (Ar C), 126.01 (Ar C), 124.76 (Ar C), 124.62 (Ar C), 119.03 (Ar C), 115.25 (Ar C), 113.49 (Ar C), 36.86 (d, JCP = 11.6 Hz, o-PCy), 31.35 (d, JCP = 17.4 Hz, ipso-PCy), 29.24 (NCMe₂), 28.67 (NCMe₂), 28.52 (d, JCP = 10.0 Hz, m-PCy), 26.40 (p-PCy), 25.05 (CHMe₂), 24.57 (d, JCP = 3.5 Hz, CHMe₂), 24.05 (d, JCP = 5.3 Hz, CHMe₂), 23.98 (CHMe₂), 21.84 (NCMe). ³¹P {¹H} NMR (121.32 MHz, C₆D₆, 298 K): δ -22.3.

UV-vis (Hexane, 25 °C) [λmax, nm (ε, mol⁻¹ L cm⁻¹)]: 253 (23,300), 439 (6,900). Anal. Calcd for C₄₄H₆₃N₂PZn: C, 73.77; H, 8.86; N, 3.91. Found: C, 73.65; H, 9.01; N, 3.87

[α-C₆H₄{C(CH₃)=N-2,6-Pr₂C₆H₃}{NH(2,6-Pr₂C₆H₃)}ZnP(C₆H₅)₂], [Ar-BDIDippZnPPh₂] (14) To a stirring solution of [Ar-BDIDippZnBr] (0.080 g, 0.134 mmol) in hexane (5 mL) a lithium diphenylphosphanide (0.026 g, 0.134 mmol) slurry in hexane (5 mL) was added. The reaction was left stirring for 36 hours where a white solid was suspended in a yellow solution. The reaction was filtered through celite and the volatiles were removed in vacuo. The resulting yellow solid was clean via ¹H NMR (0.046 g, 49%). Crystals were grown from minimum hexane. ¹H NMR (299.74 MHz, C₆D₆, 298 K) δ 7.46 (d, J = 8.5 Hz, 1H, C₆H₄), 7.29-7.03 (m, 7H, Ar H), 6.88 (m, 10H, PPh₂), 6.67 (d, J = 8.9 Hz, 1H, C₆H₄), 6.40 (t, J = 7.6 Hz, 1H, C₆H₄), 3.37 (septet, J = 6.8 Hz, 2H, CHMe₂), 2.91 (septet, J = 6.8 Hz, 2H, CHMe₂), 1.96 (s, 3H, NCMe), 1.13 (d, J = 7.3 Hz, 6H, CHMe₂), 1.10 (d, J = 7.3 Hz, 6H, CHMe₂), 1.07 (d, J = 7.0 Hz, 6H, CHMe₂),
0.95 (d, J = 6.8 Hz, 6H, CHMe2). 13C NMR (150.76 MHz, C6D6, 298 K): δ 176.04 (NCMe), 158.15 (ArC), 145.26 (d, JCP = 105.4 Hz, ipso-PPh), 139.92 (ArC), 139.67 (ArC), 139.55 (ArC), 139.88 (ArC, 135.21 (d, JCP = 18.1 Hz, m-PPh), 133.63 (d, JCP = 37.9 Hz, o-PPh), 127.17 (ArC), 126.32 (ArC), 125.56 (ArC), 125.07 (ArC), 118.78 (ArC), 115.27 (ArC), 133.63 (d, JCP = 37.9 Hz, o-PPh), 127.17 (ArC), 126.32 (ArC), 125.56 (ArC), 125.07 (ArC), 118.78 (ArC), 115.27 (ArC), 113.95 (ArC), 29.30 (CCHMe2), 28.64 (CCHMe2), 24.62 (CHMe2), 24.32 (d, JCP = 3.4 Hz, CHMe2), 23.80 (CHMe2), 23.46 (d, JCP = 5.3 Hz, CHMe2), 21.51 (NCMe).31P {1H} NMR (121.32 MHz, C6D6, 298 K): δ -47.7. UV-vis (Hexane, 25 °C) [λmax, nm (ε, mol⁻¹ L cm⁻¹)]: 254 (54,200), 436 (12,700).

[ο-C6H4(C(CH3)=N-2,6-iPr2C6H3){NH(2,6-iPr2C6H3)}Zn(S)2P(C6H11)2], [Ar-BDiPPZn(S)2PCy2]
(15) To a stirring slurry of [Ar-BDiPPZnPCy2] (0.120 g, 0.176 mmol) in toluene (5 mL) dried sulfur powder was added (0.028 g, 0.882 mmol). The reaction was left to react for 16 hours. The resulting bright yellow solution with light precipitate was filtered through celite and the volatiles were removed in vacuo. Hexane was added to the residue (2 mL) and filtered again. The resulting solution was kept at -30 °C. Bright yellow/green highly fluorescent rectangular crystals grew after 24 hours (0.087 g, 63%). 1H NMR (299.74 MHz, C6D6, 298 K): δ 7.49 (dd, J = 8.6, 1.7 Hz, 1H, C6H4), 7.40 – 7.25 (m, 6H, ArH), 6.86 (ddd, J = 8.5, 6.5, 1.7 Hz, 1H, C6H4), 6.65 (dd, J = 8.9, 1.3 Hz, 1H, C6H4), 6.35 (ddd, J = 8.4, 6.7, 1.8 Hz, 1H, C6H4), 3.65 (septet, J = 6.8 Hz, 2H, C6HMe2), 3.15 (p, J = 6.7 Hz, 2H, CHMe2), 2.09 (s, 3H, NCMe), 1.63 (d, J = 6.8 Hz, 6H, CHMe2), 1.59 (d, J = 6.8 Hz, 6H, CHMe2), 1.21 (d, J = 6.8 Hz, 6H, CHMe2), 0.96 (d, J = 6.7 Hz, 6H, CHMe2), cyclohexyl peaks were a range of multiplets from 1.55-1.13 ppm. 13C NMR (150 MHz, C6D6, 298 K): δ 175.96 (NCMe), 159.38 (ArC), 145.25 (ArC), 145.18 (ArC), 144.42 (ArC), 141.31 (ArC), 133.81 (ArC), 132.86 (ArC), 126.45 (ArC), 125.32 (ArC), 124.38 (ArC), 124.29 (ArC), 120.30 (ArC), 114.19 (ArC), 112.24 (ArC), 29.19 (CHMe2), 28.88 (CHMe2), 26.33 (CHMe2), 25.48 (br, PCy2), 25.21 (CHMe2), 24.41 (CHMe2), 24.30 (CHMe2), 23.70 (NCMe).31P {1H} NMR (121.32 MHz, C6D6, 298 K): δ 93.4. UV-vis (Hexane, 25 °C) [λmax, nm (ε, mol⁻¹ L cm⁻¹)]: 253 (61,000), 446 (13,900).

[ο-C6H4(C(CH3)=N-2,6-iPr2C6H3){NH(2,6-iPr2C6H3)}Zn(Se)2P(C6H11)2], [Ar-BDiPPZn(Se)2PCy2]
(16) To a stirring slurry of [Ar-BDiPPZnPCy2] (0.120 g, 0.176 mmol) in toluene (5 mL) dried selenium powder was added (0.070 g, 0.882 mmol). The reaction was left to react for 16 hours. The resulting orange solution with dark precipitate was filtered through celite and the volatiles were removed in vacuo. Hexane was added to the residue (2 mL) and kept at
-30 °C. Bright yellow rectangular crystals suitable for characterization and X-ray diffraction grew after 24 hours (0.062 g, 42%). \(^1\)H NMR (299.74 MHz, C\(_6\)D\(_6\), 298 K): \(\delta \) 7.46 (d, \(J = 8.7 \) Hz, 1H, C\(_6\)H\(_4\)), 7.39-7.19 (m, 6H, ArH), 6.86 (t, \(J = 8.1 \) Hz, 1H, C\(_6\)H\(_4\)), 6.66 (d, \(J = 9.5 \) Hz, 1H, C\(_6\)H\(_4\)), 6.34 (t, \(J = 7.5 \) Hz, 1H, C\(_6\)H\(_4\)), 3.67 (septet, \(J = 6.6 \) Hz, 2H, CH\(_{\text{Me}}\)_2), 3.17 (septet, \(J = 6.7 \) Hz, 2H, CH\(_{\text{Me}}\)_2), 2.09 (s, 6H, NC\(_{\text{Me}}\)), 1.64 (d, \(J = 6.8 \) Hz, 6H, CH\(_{\text{Me}}\)_2), 1.61 (d, \(J = 6.9 \) Hz, 6H, CH\(_{\text{Me}}\)_2), 1.23 (d, \(J = 6.4 \) Hz, 6H, CH\(_{\text{Me}}\)_2), 0.98 (d, \(J = 6.7 \) Hz, 6H, CH\(_{\text{Me}}\)_2), cyclohexyl peaks were a range of multiplets from 1.57-0.88 ppm. \(^{13}\)C NMR (150.76 MHz, C\(_6\)D\(_6\), 298 K): \(\delta \) 175.61 (NC\(_{\text{Me}}\)), 159.15 (ArC), 145.48 (ArC), 145.17 (ArC), 144.69 (ArC), 141.25 (ArC), 133.75 (ArC), 133.63 (ArC), 126.39 (ArC), 125.25 (ArC), 124.40 (ArC), 124.32 (ArC), 120.64 (ArC), 114.39 (ArC), 112.18 (ArC), 29.31 (CH\(_{\text{Me}}\)_2), 29.03 (CH\(_{\text{Me}}\)_2), 26.61 (PCy), 26.38 (PCy), 26.36 (d, \(J_{\text{CP}} = 15.3 \) Hz, ipso-PCy), 26.09 (PCy), 25.28 (CH\(_{\text{Me}}\)_2), 24.76 (CH\(_{\text{Me}}\)_2), 24.66 (CH\(_{\text{Me}}\)_2), 24.23 (NC\(_{\text{Me}}\)). \(^{31}\)P {\(^1\)H} NMR (121.32 MHz, C\(_6\)D\(_6\), 298 K): \(\delta \) 51.2 (J\(_{\text{PSe}}\) = 447.9 Hz). UV-vis (Hexane, 25 °C) \([\lambda_{\text{max}}, \text{nm} (\epsilon, \text{mol}^{-1} \text{L cm}^{-1})]\): 255 (58,900), 446 (10,600).

\([\text{o-C}_6\text{H}_4\{\text{C(CH}_3)\}}\text{=N-2,6-Pr}_2\text{C}_6\text{H}_3]\{\text{NH}(2,6-\text{Pr}_2\text{C}_6\text{H}_3}\}\text{Zn(Se)}_2\text{P}(\text{C}_6\text{H}_{11})_2\], [Ar-BDI\(_{\text{dipp}}\)Zn(Te)\(_2\)PCy\(_2\)]

(17) To a stirring slurry of [Ar-BDI\(_{\text{dipp}}\)ZnPCy\(_2\)] (0.030 g, 0.042 mmol) in toluene (5 mL) dried tellurium powder was added (0.011 g, 0.084 mmol). The reaction was left to react for 72 hours. The resulting orange solution with dark precipitate was filtered through celite and the volatiles were removed in vacuo. Hexane was added to the residue (2 mL) and kept at -30 °C. Orange rectangular crystals suitable for characterization and X-ray diffraction grew after 24 hours (0.017 g, 41%). \(\delta \) 7.48 – 7.21 (m, 6H, ArH), 6.93 – 6.79 (m, 2H, C\(_4\)H\(_4\)), 6.62 (d, \(J = 8.8 \) Hz, 1H, C\(_6\)H\(_4\)), 6.31 (t, \(J = 7.4 \) Hz, 1H, C\(_6\)H\(_4\)), 3.74 (septet, \(J = 6.8 \) Hz, 2H, CH\(_{\text{Me}}\)_2), 3.27 (septet, \(J = 6.7 \) Hz, 2H, CH\(_{\text{Me}}\)_2), 2.05 (s, 3H, NC\(_{\text{Me}}\)), 1.64 (d, \(J = 6.8 \) Hz, 12H, CH\(_{\text{Me}}\)_2), 1.29 (d, \(J = 6.7 \) Hz, 6H, CH\(_{\text{Me}}\)_2), 1.05 (d, \(J = 6.7 \) Hz, 6H, CH\(_{\text{Me}}\)_2). cyclohexyl peaks were a range of multiplets from 1.55-0.87 ppm. \(^{13}\)C NMR (150.76 MHz, C\(_6\)D\(_6\), 298 K): \(\delta \) 175.29 (NC\(_{\text{Me}}\)), 158.82 (ArC), 145.97 (ArC), 145.41 (ArC), 141.30 (ArC), 133.60 (ArC), 132.27 (ArC), 128.37 (ArC), 126.63 (ArC), 125.45 (ArC), 124.57 (ArC), 124.55 (ArC), 121.12 (ArC), 115.22 (ArC), 112.23 (ArC), 37.32 (d, \(J = 14.4 \) Hz, o-PCy), 29.30 (CH\(_{\text{Me}}\)_2), 29.09 (m-PCy), 28.97 (CH\(_{\text{Me}}\)_2), 26.22 (CH\(_{\text{Me}}\)_2), 26.17 (CH\(_{\text{Me}}\)_2), 26.12 (NC\(_{\text{Me}}\)_2), 26.00 (PCy), 25.86 (d, \(J = 17.4 \) Hz, ipso-PCy), 25.25 (NC\(_{\text{Me}}\)_2), 25.00 (NC\(_{\text{Me}}\)). \(^{31}\)P {\(^1\)H} NMR (121.32 MHz, C\(_6\)D\(_6\), 298 K): \(\delta \) 160.3. UV-vis (Hexane, 25 °C) \([\lambda_{\text{max}}, \text{nm} (\epsilon, \text{mol}^{-1} \text{L cm}^{-1})]\): 254 (29,700), 451 (4,400).
To a stirring slurry of [Ar-BDl_{ipp}ZnPPh_{2}] (0.090 g, 0.128 mmol) in toluene (5 mL) dried selenium powder was added (0.060 g, 0.760 mmol). The reaction was left to react for 16 hours. The resulting orange solution with dark precipitate was filtered through celite and the volatiles were removed in vacuo. Hexane was added to the residue (2 mL) and kept at -30 °C. Orange rectangular crystals grew after 24 hours but were not suitable for X-ray diffraction (0.057 g, 52%). 1H NMR (299.74 MHz, C_{6}D_{6}, 298 K): δ 7.37 (d, J = 8.7 Hz, 1H, C_{4}H_{4}), 7.34 – 7.26 (m, 5H, ArH), 6.83 (m, 12H, ArH), 6.63 (d, J = 8.9 Hz, 1H, C_{4}H_{4}), 6.33 (t, J = 7.5 Hz, 1H, C_{4}H_{4}), 3.78 (septet, J = 6.8 Hz, 2H, C_{4}H_{Me}_{2}), 3.27 (septet, J = 6.9 Hz, 2H, C_{4}H_{Me}_{2}), 2.03 (s, 3H, NC_{Me}), 1.41 (d, J = 3.6 Hz, 6H, CHMe_{2}), 1.40 (d, J = 3.5 Hz, 6H, CHMe_{2}), 1.26 (d, J = 6.8 Hz, 6H, CHMe_{2}), 1.03 (d, J = 6.7 Hz, 6H, CHMe_{2}). 13C NMR (150.76 MHz, C_{6}D_{6}, 298 K): δ 176.02 (NC_{Me}), 159.47 (ArC), 145.32 (ArC), 145.29 (ArC), 144.39 (ArC), 141.25 (ArC), 133.72 (ArC), 132.73 (ArC), 131.14 (ArC), 131.06 (ArC), 130.85 (ArC), 128.36 (ArC), 126.70 (ArC), 125.50 (ArC), 124.66 (ArC), 120.49 (ArC), 115.55 (ArC), 112.51 (ArC), 29.16 (CHMe_{2}), 28.71 (CHMe_{2}), 25.91 (CHMe_{2}), 25.31 (CHMe_{2}), 25.09 (CHMe_{2}), 24.90 (CHMe_{2}), 24.09 (NCMe). 31P {1H} NMR (121.32 MHz, C_{6}D_{6}, 298 K): δ 6.14 (J_{PSe} = 466.5 Hz). UV-vis (Hexane, 25 °C) [λ_{max}, nm (ε, mol⁻¹ L cm⁻¹)]: 252 (50,400), 446 (11,300).

To a stirring slurry of [Ar-BDl_{ipp}ZnPPh_{2}] (0.030 g, 0.043 mmol) in toluene (5 mL) dried tellurium powder was added (0.060 g, 0.470 mmol). The reaction was left to react for 48 hours. The resulting orange solution with dark precipitate was filtered through celite and the volatiles were removed in vacuo. Hexane was added to the residue (2 mL) and kept at -30 °C. Golden yellow rectangular crystals suitable for characterization and X-ray diffraction grew after 72 hours (0.037 g, 73%). 1H NMR (599.48 MHz, C_{6}D_{6}, 298 K): δ 7.34 (dd, J = 8.5, 1.6 Hz, 2H, C_{4}H_{4}), 7.28 – 7.19 (m, 18H, ArH), 7.10 – 7.00 (m, 18H, ArH), 6.81 (ddd, J = 8.6, 6.6, 1.6 Hz, 2H, C_{4}H_{4}), 6.47 (dd, J = 8.9, 1.8 Hz, 2H, C_{4}H_{4}), 6.36 (ddd, J = 8.4, 6.6, 1.5 Hz, 2H, C_{4}H_{4}), 3.40 (septet, J = 6.9 Hz, 4H, CHMe_{2}), 2.85 (septet, J = 6.8 Hz, 4H, CHMe_{2}), 1.94 (s, 6H, NCMe), 1.16 (d, J = 6.9 Hz, 12H, CHMe_{2}), 1.12 (d, J = 6.8 Hz, 12H, CHMe_{2}), 1.07 (d, J = 6.8 Hz, 12H, CHMe_{2}), 1.03 (d, J = 6.8 Hz, 12H, CHMe_{2}). 13C NMR (150.76 MHz, C_{6}D_{6}, 298 K): δ 175.16 (NCMe), 157.94 (ArC), 145.34 (ArC), 144.91 (ArC), 143.58 (ArC), 140.36 (ArC), 133.17 (ArC), 131.06 (ArC), 128.36 (ArC), 126.70 (ArC), 125.50 (ArC), 124.66 (ArC), 120.49 (ArC), 115.55 (ArC), 112.51 (ArC), 29.16 (CHMe_{2}), 28.71 (CHMe_{2}), 25.91 (CHMe_{2}), 25.31 (CHMe_{2}), 25.09 (CHMe_{2}), 24.90 (CHMe_{2}), 24.09 (NCMe). 31P {1H} NMR (121.32 MHz, C_{6}D_{6}, 298 K): δ 6.14 (J_{PSe} = 466.5 Hz). UV-vis (Hexane, 25 °C) [λ_{max}, nm (ε, mol⁻¹ L cm⁻¹)]: 252 (50,400), 446 (11,300).
132.76 (ArC), 126.86 (ArC), 125.85 (ArC), 124.83 (ArC), 124.25 (ArC), 119.33 (ArC), 117.10 (ArC), 113.79 (ArC), 29.15 (CHMe2), 28.37 (CHMe2), 25.65 (CHMe2), 25.22 (CHMe2), 24.74 (CHMe2), 24.18 (NCMe). Note the impurity [(Ph2P)2]: 1H NMR (599.48 MHz, C6D6, 298 K): δ 7.54 – 7.48 (m, 10H, PPh), 6.94 – 6.89 (m, 10H, PPh). 13C NMR (150.76 MHz, C6D6, 298 K): δ 136.61 (t, JCP = 11.1 Hz, PPh), 134.83 (t, JCP = 12.9 Hz, PPh), 128.90 (s, PPh), 128.59 (t, JCP = 7.7 Hz, PPh).
Chapter 4

Appendices

4.1 Appendix A: NMR Spectra of Novel Compounds

$^1$H NMR (299.74 MHz, C₆D₆, 298 K) and $^{31}$P NMR (121.32 MHz C₆D₆, 298 K) spectra of [BDI$_{Dipp}$ZnPCy$_2$] XXX with diethyl ether

$^{13}$C NMR spectrum of [BDI$_{Dipp}$ZnPCy$_2$] XXX (150.76 MHz, C₆D₆, 298 K) with diethyl ether
$^1$H NMR (299.74 MHz C$_6$D$_6$, 298 K) and $^{31}$P NMR (121.32 MHz C$_6$D$_6$, 298 K) spectra of [BDl$_{Dipp}$Zn(Se)$_2$PCy$_2$] XXXIII with diethyl ether

$^{13}$C NMR spectrum of [BDl$_{Dipp}$Zn(Se)$_2$PCy$_2$] XXXIII (150.76 MHz, C$_6$D$_6$, 298 K) with diethyl ether
$^{31}$P NMR (121.32 MHz, C$_6$D$_6$, 298 K) spectrum of [BDlDippZn(Te)PCy$_2$] $^{XXXV}$

$^{31}$P NMR (121.32 MHz, C$_6$D$_6$, 298 K) spectrum of [BDlDippZnPPh$_2$] $^{XXXI}$
$^1$H NMR (599.48 MHz C₆D₆, 298 K) and $^{31}$P NMR (121.32 MHz C₆D₆, 298 K) spectra of [BDlDippZn(Se)₂PPh₂] \textbf{XXXIV}

$^{13}$C NMR spectrum of [BDlDippZn(Se)₂PPh₂] \textbf{XXXIV} (150.76 MHz, C₆D₆, 298 K)
$^1$H NMR spectrum of [Ar-BDI_{Dipp}HgCl] 1 (599.48 MHz, C$_6$D$_6$, 298 K)

$^{13}$C NMR spectrum of [Ar-BDI_{Dipp}HgCl] 1 (150.76 MHz, C$_6$D$_6$, 298 K)
$^1$H NMR spectrum of [BDI$_{Dipp}$CdH] 3 (299.74 MHz, C$_6$D$_6$, 298 K)

$^{13}$C NMR spectrum of [BDI$_{Dipp}$CdH] 3 (150.76 MHz, C$_6$D$_6$, 298 K)
$^1$H NMR spectrum of [BDI$_{Dipp}$Cd]$_2$ 5 (299.74 MHz, C$_6$D$_6$, 298 K) with hexane

$^{13}$C NMR spectrum of [BDI$_{Dipp}$Cd]$_2$ 5 (150.76 MHz, C$_6$D$_6$, 298 K)
$^1$H NMR spectrum of [(DC-Am)$_2$(DC-Fo)Cd] 8 (299.74 MHz, C$_6$D$_6$, 298 K)

$^1$H NMR (299.74 MHz C$_6$D$_6$, 298 K) and $^{31}$P NMR (121.32 MHz C$_6$D$_6$, 298 K) spectra of [BDI$_{Dipp}$CdPCy$_2$] 2 with toluene and [BDI$_{Dipp}$-H] impurity
$^1$H NMR spectrum of [Ar-BDI$_{Dipp}$ZnBr$_2$Li(THF)$_4$] 9 (299.74 MHz, C$_6$D$_6$, 298 K) with toluene
$^1$H NMR spectrum of [Ar-BDI$_{Dipp}$ZnBr] $10$ (299.74 MHz, C$_6$D$_6$, 298 K) with hexane

$^{13}$C NMR spectrum of [Ar-BDI$_{Dipp}$ZnBr] $10$ (150.76 MHz, C$_6$D$_6$, 298 K) with hexane
\(^1\)H NMR spectrum of [Ar-BDI\textsubscript{Dipp}ZnHMDS] \textbf{11} (599.48 MHz, C\textsubscript{6}D\textsubscript{6}, 298 K)

\[^{13}\text{C} \) NMR spectrum of [Ar-BDI\textsubscript{Dipp}ZnHMDS] \textbf{11} (150.76 MHz, C\textsubscript{6}D\textsubscript{6}, 298 K)
$^1$H NMR spectrum of [Ar-BDI_{Dipp}ZnNH(Dipp)] \textbf{12} (599.48 MHz, C$_6$D$_6$, 298 K)

$^{13}$C NMR spectrum of [Ar-BDI_{Dipp}ZnNH(Dipp)] \textbf{12} (150.76 MHz, C$_6$D$_6$, 298 K) with hexane
$^1$H NMR (599.48 MHz) and $^{31}$P NMR (121.32 MHz C₆D₆, 298 K) spectra of [Ar-BDIdippZnPCy₂] 13 with hexane

$^{13}$C NMR spectrum of [Ar-BDIdippZnPCy₂] 13 (150.76 MHz, C₆D₆, 298 K) with hexane
$^1$H NMR (299.74 MHz, C$_6$D$_6$, 298 K) and $^{31}$P NMR (121.32 MHz, C$_6$D$_6$, 298 K) spectra of [Ar-BDI$_{Dipp}$Zn(S)$_2$PCy$_2$] 15 with hexane

$^{13}$C NMR spectrum of [Ar-BDI$_{Dipp}$Zn(S)$_2$PCy$_2$] 15 (150.76 MHz, C$_6$D$_6$, 298 K) with hexane
$^1$H NMR (599.48 MHz, C$_6$D$_6$, 298 K) and $^{31}$P NMR (121.32 MHz, C$_6$D$_6$, 298 K) spectra of [Ar-BDI$_{Dipp}$Zn(Se)$_2$PCy$_2$] 16 with unreacted [Ar-BDI$_{Dipp}$ZnPCy$_2$]

$^{13}$C NMR spectrum of [Ar-BDI$_{Dipp}$Zn(Se)$_2$PCy$_2$] 16 (150.76 MHz, C$_6$D$_6$, 298 K) with hexane
$^1$H NMR (599.48 MHz) and $^{31}$P NMR (121.32 MHz, C$_6$D$_6$, 298 K) spectra of [Ar-BDl$_{Dipp}$Zn( Te)$_2$PCy$_2$] 17

$^{13}$C NMR spectrum of [Ar-BDl$_{Dipp}$Zn( Te)$_2$PCy$_2$] 17 (150.74 MHz, C$_6$D$_6$, 298 K) with hexane
$^1$H NMR (599.48 MHz, C$_6$D$_6$, 298 K) and $^{31}$P NMR (121.32 MHz, C$_6$D$_6$, 298 K) spectra of [Ar-BDlDippZnPPh$_2$] 14 with unreacted [Ar-BDlDippZnBr]

$^{13}$C NMR spectrum of [Ar-BDlDippZnPPh$_2$] 14 (150.74 MHz, C$_6$D$_6$, 298 K)
$^1$H NMR (299.74 MHz, C$_6$D$_6$, 298 K) and $^{31}$P NMR (121.32 MHz, C$_6$D$_6$, 298 K) spectra of [Ar-BDl$_{Dipp}$Zn(Se)$_2$PPh$_2$] 18

$^{13}$C NMR spectrum of [Ar-BDl$_{Dipp}$Zn(Se)$_2$PCy$_2$] 18 (150.76 MHz, C$_6$D$_6$, 298 K) with hexane
$^1$H NMR spectrum of [Ar-BDI$_{Dipp}$Zn]$_2$Te 19 (599.48 MHz, C$_6$D$_6$, 298 K) with hexane and (Ph$_2$P)$_2$

$^{13}$C NMR spectrum of [Ar-BDI$_{Dipp}$Zn]$_2$Te 19 (150.76 MHz, C$_6$D$_6$, 298 K) with hexane and [(Ph$_2$P)$_2$]
4.2 Appendix B: Crystal Data Tables

Crystal data and structure refinement for 55rf12. (XXX)

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<th>Parameter</th>
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<td>Temperature/K</td>
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<td>Crystal system</td>
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<td>P2₁</td>
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<td>a/Å</td>
<td>11.11426(14)</td>
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<td>b/Å</td>
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</tr>
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<td>13.06175(18)</td>
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<td>ρcalc g/cm³</td>
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<td>F(000)</td>
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<td>Crystal size/mm³</td>
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<tr>
<td>Radiation</td>
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<td>2θ range for data collection/°</td>
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<td>Reflections collected</td>
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<td>Goodness-of-fit on F²</td>
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<tr>
<td>Final R indexes [I&gt;=2σ (I)]</td>
<td>R₁ = 0.0481, wR₂ = 0.1174</td>
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<td>Final R indexes [all data]</td>
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<tr>
<td>Flack parameter</td>
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The crystal structure has disorder in the phosphanide and the cyclohexyl groups which lay in two directions, 81% in one direction and 19% in the other. This was modelled, although the carbon atoms of the lower occupancy component were left isotropic.
Crystal data and structure refinement for 55RF13. (XXXII)

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<td>Temperature/K</td>
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<td>c/Å</td>
<td>15.7842(4)</td>
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<tr>
<td>β/°</td>
<td>99.159(3)</td>
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<td>γ/°</td>
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<td>Volume/Å³</td>
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<td>Crystal size/mm³</td>
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<td>2Θ range for data collection/°</td>
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<td>Crystal data and structure refinement for 55RF11. (XXXIII)</td>
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<td>10.40514(14)</td>
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<td>( \rho_{\text{calc}}/\text{g/cm}³ )</td>
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<td>( \mu/\text{mm}⁻¹ )</td>
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<td>Crystal size/mm³</td>
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Crystal data and structure refinement for 55rf10. (XXXIV)

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<td>Space group</td>
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<td>11.5002(3)</td>
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<td>γ/°</td>
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<td>MoKα ($\lambda = 0.71073$)</td>
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<td>5.688 to 70.548</td>
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<td>$-19 \leq h \leq 19$, $-25 \leq k \leq 24$, $-21 \leq l \leq 22$</td>
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</tr>
<tr>
<td><strong>Independent reflections</strong></td>
<td>11367 [R$<em>{\text{int}} = 0.0347$, R$</em>{\text{sigma}} = 0.0389$]</td>
</tr>
<tr>
<td><strong>Data/restraints/parameters</strong></td>
<td>11367/0/303</td>
</tr>
<tr>
<td><strong>Goodness-of-fit on F$^2$</strong></td>
<td>1.079</td>
</tr>
<tr>
<td><strong>Final R indexes [I&gt;=2σ (I)]</strong></td>
<td>R$_1$ = 0.0330, wR$_2$ = 0.0772</td>
</tr>
<tr>
<td><strong>Final R indexes [all data]</strong></td>
<td>R$_1$ = 0.0489, wR$_2$ = 0.0874</td>
</tr>
<tr>
<td><strong>Largest diff. peak/hole / e Å$^{-3}$</strong></td>
<td>1.17/-0.81</td>
</tr>
<tr>
<td>Crystal data and structure refinement for 55rf21. (4)</td>
<td></td>
</tr>
<tr>
<td>-----------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td><strong>Identification code</strong></td>
<td>55rf21</td>
</tr>
<tr>
<td><strong>Empirical formula</strong></td>
<td>C₅₈H₈₂Cd₂N₄</td>
</tr>
<tr>
<td><strong>Formula weight</strong></td>
<td>1060.07</td>
</tr>
<tr>
<td><strong>Temperature/K</strong></td>
<td>278.07(10)</td>
</tr>
<tr>
<td><strong>Crystal system</strong></td>
<td>monoclinic</td>
</tr>
<tr>
<td><strong>Space group</strong></td>
<td>P2₁/c</td>
</tr>
<tr>
<td><strong>a/Å</strong></td>
<td>14.4118(4)</td>
</tr>
<tr>
<td><strong>b/Å</strong></td>
<td>14.2417(4)</td>
</tr>
<tr>
<td><strong>c/Å</strong></td>
<td>26.8111(8)</td>
</tr>
<tr>
<td><strong>α/°</strong></td>
<td>90</td>
</tr>
<tr>
<td><strong>β/°</strong></td>
<td>94.817(3)</td>
</tr>
<tr>
<td><strong>γ/°</strong></td>
<td>90</td>
</tr>
<tr>
<td><strong>Volume/Å³</strong></td>
<td>5483.5(3)</td>
</tr>
<tr>
<td><strong>Z</strong></td>
<td>4</td>
</tr>
<tr>
<td><strong>ρ&lt;sub&gt;calc&lt;/sub&gt;/cm³</strong></td>
<td>1.284</td>
</tr>
<tr>
<td><strong>μ/mm⁻¹</strong></td>
<td>0.814</td>
</tr>
<tr>
<td><strong>F(000)</strong></td>
<td>2216.0</td>
</tr>
<tr>
<td><strong>Crystal size/mm³</strong></td>
<td>0.20 × 0.21 × 0.05</td>
</tr>
<tr>
<td><strong>Radiation</strong></td>
<td>MoKα (λ = 0.71073)</td>
</tr>
<tr>
<td><strong>2θ range for data collection/°</strong></td>
<td>5.396 to 54.998</td>
</tr>
<tr>
<td><strong>Index ranges</strong></td>
<td>-18 ≤ h ≤ 18, -18 ≤ k ≤ 18, -34 ≤ l ≤ 34</td>
</tr>
<tr>
<td><strong>Reflections collected</strong></td>
<td>60496</td>
</tr>
<tr>
<td><strong>Independent reflections</strong></td>
<td>12588 [R&lt;sub&gt;int&lt;/sub&gt; = 0.0470, R&lt;sub&gt;sigma&lt;/sub&gt; = 0.0364]</td>
</tr>
<tr>
<td><strong>Data/restraints/parameters</strong></td>
<td>12588/0/597</td>
</tr>
<tr>
<td><strong>Goodness-of-fit on F²</strong></td>
<td>1.117</td>
</tr>
<tr>
<td><strong>Final R indexes [I&gt;2σ (I)]</strong></td>
<td>R&lt;sub&gt;1&lt;/sub&gt; = 0.0352, wR&lt;sub&gt;2&lt;/sub&gt; = 0.0746</td>
</tr>
<tr>
<td><strong>Final R indexes [all data]</strong></td>
<td>R&lt;sub&gt;1&lt;/sub&gt; = 0.0471, wR&lt;sub&gt;2&lt;/sub&gt; = 0.0812</td>
</tr>
<tr>
<td><strong>Largest diff. peak/hole / e Å⁻³</strong></td>
<td>0.89/-0.71</td>
</tr>
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Crystal data and structure refinement for exp_82. (10)

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<th>Identification code</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Empirical formula</td>
<td>C_{32}H_{41}BrN_{2}Zn</td>
</tr>
<tr>
<td>Formula weight</td>
<td>598.95</td>
</tr>
<tr>
<td>Temperature/K</td>
<td>291.24(10)</td>
</tr>
<tr>
<td>Crystal system</td>
<td>trigonal</td>
</tr>
<tr>
<td>Space group</td>
<td>R-3</td>
</tr>
<tr>
<td>a/Å</td>
<td>34.4281(6)</td>
</tr>
<tr>
<td>b/Å</td>
<td>34.4281(6)</td>
</tr>
<tr>
<td>c/Å</td>
<td>14.1576(3)</td>
</tr>
<tr>
<td>α/°</td>
<td>90</td>
</tr>
<tr>
<td>β/°</td>
<td>90</td>
</tr>
<tr>
<td>γ/°</td>
<td>120</td>
</tr>
<tr>
<td>Volume/Å³</td>
<td>14532.6(6)</td>
</tr>
<tr>
<td>Z</td>
<td>18</td>
</tr>
<tr>
<td>ρ calc g/cm³</td>
<td>1.232</td>
</tr>
<tr>
<td>μ/mm⁻¹</td>
<td>2.643</td>
</tr>
<tr>
<td>F(000)</td>
<td>5616.0</td>
</tr>
<tr>
<td>Crystal size/mm³</td>
<td>0.2868 × 0.2433 × 0.1644</td>
</tr>
<tr>
<td>Radiation</td>
<td>CuKα (λ = 1.54184)</td>
</tr>
<tr>
<td>2Θ range for data collection/°</td>
<td>8.614 to 143.53</td>
</tr>
<tr>
<td>Index ranges</td>
<td>-41 ≤ h ≤ 42, -31 ≤ k ≤ 38, -17 ≤ l ≤ 16</td>
</tr>
<tr>
<td>Reflections collected</td>
<td>16028</td>
</tr>
<tr>
<td>Independent reflections</td>
<td>6236 [R_{int} = 0.0214, R_{sigma} = 0.0235]</td>
</tr>
<tr>
<td>Data/restraints/parameters</td>
<td>6236/0/356</td>
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<tr>
<td>Goodness-of-fit on F²</td>
<td>1.050</td>
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<tr>
<td>Final R indexes [I&gt;=2σ (I)]</td>
<td>R₁ = 0.0249, wR₂ = 0.0613</td>
</tr>
<tr>
<td>Final R indexes [all data]</td>
<td>R₁ = 0.0270, wR₂ = 0.0624</td>
</tr>
<tr>
<td>Largest diff. peak/hole / e Å⁻³</td>
<td>0.31/-0.28</td>
</tr>
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</table>

The crystal structure has disorder in the BDI backbone which lay in two directions, 75% in one direction and 25% in the other. This was modelled, although the carbon atoms of the lower occupancy component were left isotropic.
<table>
<thead>
<tr>
<th>Identification code</th>
<th>exp_75</th>
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</thead>
<tbody>
<tr>
<td>Empirical formula</td>
<td>C_{76}H_{118}N_{6}Si_{4}Zn_{2}</td>
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<tr>
<td>Formula weight</td>
<td>1358.86</td>
</tr>
<tr>
<td>Temperature/K</td>
<td>120.01(10)</td>
</tr>
<tr>
<td>Crystal system</td>
<td>orthorhombic</td>
</tr>
<tr>
<td>Space group</td>
<td>Pca2_1</td>
</tr>
<tr>
<td>a/Å</td>
<td>18.1072(2)</td>
</tr>
<tr>
<td>b/Å</td>
<td>19.7437(3)</td>
</tr>
<tr>
<td>c/Å</td>
<td>21.5080(3)</td>
</tr>
<tr>
<td>α/°</td>
<td>90</td>
</tr>
<tr>
<td>β/°</td>
<td>90</td>
</tr>
<tr>
<td>γ/°</td>
<td>90</td>
</tr>
<tr>
<td>Volume/Å^3</td>
<td>7689.20(17)</td>
</tr>
<tr>
<td>Z</td>
<td>4</td>
</tr>
<tr>
<td>ρ_{calc}/g/cm^3</td>
<td>1.174</td>
</tr>
<tr>
<td>μ/mm^{-1}</td>
<td>1.686</td>
</tr>
<tr>
<td>F(000)</td>
<td>2928.0</td>
</tr>
<tr>
<td>Crystal size/mm^3</td>
<td>0.1931 x 0.1713 x 0.0946</td>
</tr>
<tr>
<td>Radiation</td>
<td>CuKα (λ = 1.54184)</td>
</tr>
<tr>
<td>2θ range for data collection/°</td>
<td>9.364 to 143.538</td>
</tr>
<tr>
<td>Index ranges</td>
<td>-22 ≤ h ≤ 20, -24 ≤ k ≤ 19, -26 ≤ l ≤ 26</td>
</tr>
<tr>
<td>Reflections collected</td>
<td>34922</td>
</tr>
<tr>
<td>Independent reflections</td>
<td>13699 [R_{int} = 0.0205, R_{sigma} = 0.0237]</td>
</tr>
<tr>
<td>Data/restraints/parameters</td>
<td>13699/1/803</td>
</tr>
<tr>
<td>Goodness-of-fit on F^2</td>
<td>1.033</td>
</tr>
<tr>
<td>Final R indexes [I&gt;=2σ (I)]</td>
<td>R_1 = 0.0329, wR_2 = 0.0833</td>
</tr>
<tr>
<td>Final R indexes [all data]</td>
<td>R_1 = 0.0344, wR_2 = 0.0846</td>
</tr>
<tr>
<td>Largest diff. peak/hole / e Å^{-3}</td>
<td>0.94/-0.97</td>
</tr>
<tr>
<td>Flack parameter</td>
<td>-0.007(7)</td>
</tr>
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Table 1 Crystal data and structure refinement for exp_79. (13)

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<tr>
<th>Parameter</th>
<th>Value/Condition</th>
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<tbody>
<tr>
<td>Identification code</td>
<td>exp_79</td>
</tr>
<tr>
<td>Empirical formula</td>
<td>C_{44}H_{63}N_{2}PZn</td>
</tr>
<tr>
<td>Formula weight</td>
<td>716.3</td>
</tr>
<tr>
<td>Temperature/K</td>
<td>120.01(10)</td>
</tr>
<tr>
<td>Crystal system</td>
<td>monoclinic</td>
</tr>
<tr>
<td>Space group</td>
<td>P2\textsubscript{1}/n</td>
</tr>
<tr>
<td>a/Å</td>
<td>10.88553(12)</td>
</tr>
<tr>
<td>b/Å</td>
<td>16.51628(19)</td>
</tr>
<tr>
<td>c/Å</td>
<td>22.0120(2)</td>
</tr>
<tr>
<td>α/°</td>
<td>90</td>
</tr>
<tr>
<td>β/°</td>
<td>91.6596(10)</td>
</tr>
<tr>
<td>γ/°</td>
<td>90</td>
</tr>
<tr>
<td>Volume/Å\textsuperscript{3}</td>
<td>3955.84(8)</td>
</tr>
<tr>
<td>Z</td>
<td>4</td>
</tr>
<tr>
<td>ρ calcg/cm\textsuperscript{3}</td>
<td>1.203</td>
</tr>
<tr>
<td>μ/mm\textsuperscript{-1}</td>
<td>1.464</td>
</tr>
<tr>
<td>F(000)</td>
<td>1544.0</td>
</tr>
<tr>
<td>Crystal size/mm\textsuperscript{3}</td>
<td>0.2118 × 0.1523 × 0.0757</td>
</tr>
<tr>
<td>Radiation</td>
<td>Cu Kα (λ = 1.54184)</td>
</tr>
<tr>
<td>2Θ range for data collection/°</td>
<td>8.9616 to 143.6478</td>
</tr>
<tr>
<td>Index ranges</td>
<td>-13 ≤ h ≤ 13, -19 ≤ k ≤ 20, -27 ≤ l ≤ 27</td>
</tr>
<tr>
<td>Reflections collected</td>
<td>28122</td>
</tr>
<tr>
<td>Independent reflections</td>
<td>7730 [R_{int} = 0.0334, R_{sigma} = 0.0407]</td>
</tr>
<tr>
<td>Data/restraints/parameters</td>
<td>7730/0/482</td>
</tr>
<tr>
<td>Goodness-of-fit on F\textsuperscript{2}</td>
<td>1.099</td>
</tr>
<tr>
<td>Final R indexes [I≥2σ (I)]</td>
<td>R_1 = 0.0611, wR_2 = 0.1593</td>
</tr>
<tr>
<td>Final R indexes [all data]</td>
<td>R_1 = 0.0658, wR_2 = 0.1632</td>
</tr>
<tr>
<td>Largest diff. peak/hole / e Å\textsuperscript{-3}</td>
<td>1.23/-0.70</td>
</tr>
</tbody>
</table>

The crystal structure has disorder in one of the cyclohexyl groups which lay in two directions, 40% in one direction and 60% in the other. The crystal structure has disorder in one of the isopropyl groups which lay in two directions, 42% in one direction and 58% in the other. Both were modelled, although the carbon atoms of the lower occupancy component were left isotropic.
Crystal data and structure refinement for exp_84. (14)

<table>
<thead>
<tr>
<th>Identification code</th>
<th>exp_84</th>
</tr>
</thead>
<tbody>
<tr>
<td>Empirical formula</td>
<td>C_{44}H_{51}N_{2}PZn</td>
</tr>
<tr>
<td>Formula weight</td>
<td>704.20</td>
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<tr>
<td>Temperature/K</td>
<td>291.24(10)</td>
</tr>
<tr>
<td>Crystal system</td>
<td>triclinic</td>
</tr>
<tr>
<td>Space group</td>
<td>P-1</td>
</tr>
<tr>
<td>a/Å</td>
<td>10.9460(3)</td>
</tr>
<tr>
<td>b/Å</td>
<td>13.4527(3)</td>
</tr>
<tr>
<td>c/Å</td>
<td>13.6087(4)</td>
</tr>
<tr>
<td>α/°</td>
<td>97.5890(19)</td>
</tr>
<tr>
<td>β/°</td>
<td>104.346(2)</td>
</tr>
<tr>
<td>γ/°</td>
<td>98.2482(19)</td>
</tr>
<tr>
<td>Volume/Å³</td>
<td>1891.77(8)</td>
</tr>
<tr>
<td>Z</td>
<td>2</td>
</tr>
<tr>
<td>ρ\text{calc}/cm³</td>
<td>1.236</td>
</tr>
<tr>
<td>μ/mm⁻¹</td>
<td>1.533</td>
</tr>
<tr>
<td>F(000)</td>
<td>748.0</td>
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<tr>
<td>Crystal size/mm³</td>
<td>0.1862 × 0.1787 × 0.1364</td>
</tr>
<tr>
<td>Radiation</td>
<td>CuKα (λ = 1.54184)</td>
</tr>
<tr>
<td>2Θ range for data collection/°</td>
<td>8.708 to 143.41</td>
</tr>
<tr>
<td>Index ranges</td>
<td>-13 ≤ h ≤ 13, -13 ≤ k ≤ 16, -16 ≤ l ≤ 14</td>
</tr>
<tr>
<td>Reflections collected</td>
<td>12976</td>
</tr>
<tr>
<td>Independent reflections</td>
<td>7215 [R_{int} = 0.0182, R_{sigma} = 0.0283]</td>
</tr>
<tr>
<td>Data/restraints/parameters</td>
<td>7215/7/442</td>
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<tr>
<td>Goodness-of-fit on F²</td>
<td>1.041</td>
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<tr>
<td>Final R indexes [l&gt;2\sigma (l)]</td>
<td>R₁ = 0.0319, wR₂ = 0.0819</td>
</tr>
<tr>
<td>Final R indexes [all data]</td>
<td>R₁ = 0.0342, wR₂ = 0.0836</td>
</tr>
<tr>
<td>Largest diff. peak/hole / e Å⁻³</td>
<td>0.40/-0.29</td>
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</tbody>
</table>
Crystal data and structure refinement for exp_86. (16)

<table>
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<th>exp_86</th>
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<td>Empirical formula</td>
<td>C₄₇H₇₀N₂PSe₂Zn</td>
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<tr>
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</tr>
<tr>
<td>Temperature/K</td>
<td>120.01(10)</td>
</tr>
<tr>
<td>Crystal system</td>
<td>monoclinic</td>
</tr>
<tr>
<td>Space group</td>
<td>P2₁/n</td>
</tr>
<tr>
<td>a/Å</td>
<td>19.06601(16)</td>
</tr>
<tr>
<td>b/Å</td>
<td>10.35149(8)</td>
</tr>
<tr>
<td>c/Å</td>
<td>24.6114(3)</td>
</tr>
<tr>
<td>α/°</td>
<td>90</td>
</tr>
<tr>
<td>β/°</td>
<td>112.7016(11)</td>
</tr>
<tr>
<td>γ/°</td>
<td>90</td>
</tr>
<tr>
<td>Volume/Å³</td>
<td>4481.04(8)</td>
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<td>Z</td>
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<td>ρ calcg/cm³</td>
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<td>3.195</td>
</tr>
<tr>
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<td>1916.0</td>
</tr>
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<td>Crystal size/mm³</td>
<td>0.2631 × 0.1445 × 0.1012</td>
</tr>
<tr>
<td>Radiation</td>
<td>CuKα (λ = 1.54184)</td>
</tr>
<tr>
<td>2Θ range for data collection/°</td>
<td>9.914 to 143.32</td>
</tr>
<tr>
<td>Index ranges</td>
<td>-14 ≤ h ≤ 23, -12 ≤ k ≤ 12, -30 ≤ l ≤ 25</td>
</tr>
<tr>
<td>Reflections collected</td>
<td>31720</td>
</tr>
<tr>
<td>Independent reflections</td>
<td>8690 [Rint = 0.0133, Rsigma = 0.0104]</td>
</tr>
<tr>
<td>Data/restraints/parameters</td>
<td>8690/3/473</td>
</tr>
<tr>
<td>Goodness-of-fit on F²</td>
<td>1.147</td>
</tr>
<tr>
<td>Final R indexes [I&gt;=2σ (I)]</td>
<td>R₁ = 0.0423, wR₂ = 0.1282</td>
</tr>
<tr>
<td>Final R indexes [all data]</td>
<td>R₁ = 0.0430, wR₂ = 0.1287</td>
</tr>
<tr>
<td>Largest diff. peak/hole / e Å⁻³</td>
<td>2.08/-0.46</td>
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<tr>
<td>Property</td>
<td>Value</td>
</tr>
<tr>
<td>--------------------------------------</td>
<td>------------------------------</td>
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<td>exp_90</td>
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<tr>
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<td>971.50</td>
</tr>
<tr>
<td>Temperature/K</td>
<td>120.01(10)</td>
</tr>
<tr>
<td>Crystal system</td>
<td>monoclinic</td>
</tr>
<tr>
<td>Space group</td>
<td>P2_{1}/n</td>
</tr>
<tr>
<td>a/Å</td>
<td>19.1598(6)</td>
</tr>
<tr>
<td>b/Å</td>
<td>10.5401(3)</td>
</tr>
<tr>
<td>c/Å</td>
<td>24.5244(9)</td>
</tr>
<tr>
<td>α/°</td>
<td>90</td>
</tr>
<tr>
<td>β/°</td>
<td>112.768(4)</td>
</tr>
<tr>
<td>γ/°</td>
<td>90</td>
</tr>
<tr>
<td>Volume/Å³</td>
<td>4566.7(3)</td>
</tr>
<tr>
<td>Z</td>
<td>10</td>
</tr>
<tr>
<td>ρ calc/g/cm³</td>
<td>3.533</td>
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<td>μ/mm⁻¹</td>
<td>4.637</td>
</tr>
<tr>
<td>F(000)</td>
<td>4900.0</td>
</tr>
<tr>
<td>Crystal size/mm³</td>
<td>0.2506 × 0.2012 × 0.0946</td>
</tr>
<tr>
<td>Radiation</td>
<td>MoKα (λ = 0.71073)</td>
</tr>
<tr>
<td>2θ range for data collection/°</td>
<td>6.646 to 59.032</td>
</tr>
<tr>
<td>Index ranges</td>
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<tr>
<td>Reflections collected</td>
<td>24889</td>
</tr>
<tr>
<td>Independent reflections</td>
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<tr>
<td>Goodness-of-fit on F²</td>
<td>1.134</td>
</tr>
<tr>
<td>Final R indexes [I&gt;=2σ (I)]</td>
<td>R₁ = 0.0466, wR₂ = 0.1568</td>
</tr>
<tr>
<td>Final R indexes [all data]</td>
<td>R₁ = 0.0589, wR₂ = 0.1677</td>
</tr>
<tr>
<td>Largest diff. peak/hole / e Å⁻³</td>
<td>2.26/-0.87</td>
</tr>
<tr>
<td>Identification code</td>
<td>exp_94b</td>
</tr>
<tr>
<td>---------------------</td>
<td>--------</td>
</tr>
<tr>
<td>Empirical formula</td>
<td>C_{67}H_{89}N_{4}TeZn_{2}</td>
</tr>
<tr>
<td>Formula weight</td>
<td>1208.76</td>
</tr>
<tr>
<td>Temperature/K</td>
<td>120.00(10)</td>
</tr>
<tr>
<td>Crystal system</td>
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</tr>
<tr>
<td>Space group</td>
<td>P-1</td>
</tr>
<tr>
<td>a/Å</td>
<td>12.1052(5)</td>
</tr>
<tr>
<td>b/Å</td>
<td>12.8767(5)</td>
</tr>
<tr>
<td>c/Å</td>
<td>22.5863(11)</td>
</tr>
<tr>
<td>α/°</td>
<td>85.696(3)</td>
</tr>
<tr>
<td>β/°</td>
<td>78.852(4)</td>
</tr>
<tr>
<td>γ/°</td>
<td>63.031(4)</td>
</tr>
<tr>
<td>Volume/Å³</td>
<td>3078.3(2)</td>
</tr>
<tr>
<td>Z</td>
<td>2</td>
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<tr>
<td>ρ_{calc}/g/cm³</td>
<td>1.304</td>
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<tr>
<td>μ/mm⁻¹</td>
<td>4.928</td>
</tr>
<tr>
<td>F(000)</td>
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<tr>
<td>Crystal size/mm³</td>
<td>0.1726 × 0.112 × 0.0541</td>
</tr>
<tr>
<td>Radiation</td>
<td>CuKα (λ = 1.54184)</td>
</tr>
<tr>
<td>2Θ range for data collection/°</td>
<td>8.33 to 144.242</td>
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<td>Index ranges</td>
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<tr>
<td>Largest diff. peak/hole / e Å⁻³</td>
<td>1.40/-1.19</td>
</tr>
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</table>
4.3 Appendix C: Bibliography


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