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SYNTHESIS, PROPERTIES, STRUCTURAL CHARACTERIZATION, AND REACTIVITY OF LOW-VALENT TITANIUM (BISDIIMINE) COMPLEXES

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ABSTRACT OF THESIS

SYNTHESIS, PROPERTIES, STRUCTURAL CHARACTERIZATION, AND REACTIVITY OF LOW-VALENT TITANIUM (BISDIIMINE) COMPLEXES

The synthesis, structure, and reactivity of titanium bis(diimine) complexes supported by 1,2-alternate dimethylsilyl-bridged p-tert-butylcalix[4]arene dianion and 2,2' methylene-bridged 4-methyl, 6-tertbutyl phenol ligands is reported. The molecular structure of [(DMSC)Ti(bpy)2] (**28**) and [(MBMP)Ti(bpy)2] (**55**) was characterized by X-ray crystallography. Complexes [(DMSC)Ti(bpy)2] (**28**), [(DMSC)Ti(dmbpy)2] (**29**), and [(DMSC)Ti(phen)2] (**30**) undergoes light-assisted reactions with two or more equivalents of (C6H5)2CO or (p-MeC6H4)2CO to give the corresponding 1-aza-5-oxatitanacyclopentene complexes **37**-**42**. Similar reactivity was observed with [(MBMP)Ti(bpy)2] (**55**), [(MBMP)Ti(dmbpy)2] (**56**), and [(MBMP)Ti(phen)2] (**57**). The molecular structure of $[(MBMP)T_i\{kappa-3-OC(C6H5)2C10H7N2\}OCH(C6H5)2]\]$ (**58**) was characterized by 1H and 13C NMR as well as X-Ray crystallography.

KEYWORDS: low-valent titanium, aromaticity, diimine complexes, X-ray crystallography, fluxionality

Marc Steven Maynor

July 28, 2004

Marc S. Maynor 2004

SYNTHESIS, PROPERTIES, STRUCTURAL CHARACTERIZATION, AND REACTIVITY OF LOW-VALENT TITANIUM BIS (DIIMINE) COMPLEXES

By

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THESIS

Marc Steven Maynor

The Graduate School

University of Kentucky

2004

SYNTHESIS, PROPERTIES, STRUCTURAL CHARACTERIZATION, AND REACTIVITY OF TITANIUM (BISDIIMINE) COMPLEXES

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A thesis submitted in partial fulfillment of the requirements for the degree of Master of Science in the College of Arts and Sciences at the University of Kentucky

By

Marc Steven Maynor

Lexington, Kentucky

Director: Dr. Folami T. Ladipo, Associate Professor of Chemistry

Lexington, Kentucky

2004

This is dedicated to all of my family and friends

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iii

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TABLE OF CONTENTS

LIST OF TABLES

LIST OF FIGURES

LIST OF SCHEMES

List of files

Chapter One

Intoduction

1.1 Intent of Thesis

The intent of this research was to synthesize and structurally characterize low-valent titanium bis (diimine) complexes. The properties and reactivity of these complexes with organic substrates were investigated. Low-valent titanium (LVT) species have been shown to have a wide range of applications in organic syntheses, such as McMurry chemistry, $1-7$ substitution of nitrobenzenes, δ cyclodimerization of unsaturated ketones⁹, coupling of chiral diimines, δ ¹⁰ oligomerization of olefins, acetylenes, and carbonyl derivatives, 11 and reduction of nitrobenzenes to amino arenes.^{12,13} Commonly, LVT species used in organic synthesis are generated by reduction of titanium tetrachloride or titanium trichloride or Ti (IV) complexes of the type L_2TiX_2 (L = Cp, NR₂, and X = Cl, Br, or OR) with common electropositive reducing agents such as Na, Zn, Li, Mg, and K.

There are a few drawbacks to the "black box" approach in organic synthesis. The LVT species is usually generated *in situ* leading to ambiguity regarding (i) the oxidation state of the titanium, (ii) the reactive species in the reaction, as well as (iii) the mechanism of the reaction. Understanding the mechanism and the properties of the reactive titanium species is important in order to expand the use of LVT. Therefore, there is great interest in synthesis and reactivity of well-characterized LVT complexes.

1.2 Background

The commercially available $TiCl₂ (1)$ and $TiCl₃ (2)$ are two of the simplest examples of LVT species. Walton and coworkers have found that **1** will react with MeCN to form [TiCl2(MeCN)] (**3).** The reaction of **3** with various donor molecules results in compounds of the type $[TiCl₂L₂]$ or $[TiCl₂B]$, $(L =$ mondentate ligand pyridine, tetrahydrofuran, or tetrahydropyran; $B =$ bidentate ligand 2,2'-bipyridyl or 1,10-phenanthroline). In these cases the Ti was assigned a +2 oxidation state.¹⁴ A series of TiE₂ (E = Cl, F, OBuⁿ, Oprⁱ and 0.5 NPh- $CH₂-CH₂-NPh$) compounds was prepared by Eisch and coworkers by the alkylative reduction of TiE₄ with two equivalents of *n*-butyllithium.¹⁵ (eq. 1.1)

TiE_4 + **2BuLi** $\overrightarrow{THF/Toluene}$ $Ti(E)_2$ + **2LiE** (eq. 1.1) **-78 C**

Without question, the use of metallocenes $[(\eta^5-C_5R_5)_2M]$ has been a milestone in the development of organometallic chemistry. Scheme 1.1 depicts a few examples of stable, wellcharacterized Ti (II) complexes supported by cyclopentadienyl ligands. Bis(η^5 cyclopentadienyl)titanium (II) dicarbonyl (4) was prepared by Bercaw and coworkers¹⁶ (Scheme 1.1). The authors found the compound to be diamagnetic. Hückel molecular orbital calculations were carried out for this complex showing that there is back-donation of electron density from the filled metal orbitals into the π^* -orbitals of the carbonyl ligands. These calculations indicated that this back-bonding is important for stabilizing the electron-rich Ti (II) center.¹⁷ Reacting 4 with 2,2-bipyridyl led to the formation of bis(η^5 -cyclopentadienyl)titanium 2,2'-bipyridyl **(5**)¹⁸ (Scheme 1.1).

Calderazzo and coworkers were able to prepare **5** by reacting $\text{bis}(\eta^5\text{-cyclopentadienyl})$ titanium dichloride with the dilithium salt of 2,2-bipyridine dianion.¹⁹ 5 was found to be weakly paramagnetic giving a room temperature EPR signal.^{18,19} McPherson and coworkers conducted magnetic susceptibility and EPR studies on **5**. The results of this study suggest that the system is diamagnetic in the ground state but has a paramagnetic exited state that is thermally accessible. The authors concluded that one unpaired electron formally occupies a molecular orbital which is localized on bis(η^5 -cyclopentadienyl) titanium while the second unpaired electron resides in the lowest energy anti-bonding orbital of the bipyridyl group.

Lawless and coworkers were able to isolate the $\frac{\text{bis}}{\text{cis}(\eta^5 \cdot \text{(tert-butyldimethylsilyl)}\cdot \text{trikl})}$ tetramethylcyclopentadienyl)] titanium (II) complex (6) by the reduction of $[\{\eta^5-C_5Me_4(SiMe_2$ t -Bu) $\{7T^2$ with sodium amalgam in toluene (Scheme 1.1). Following the discovery by Lawless, Mach and co workers were able to isolate and characterize the analogous $\{\eta^5\}$ $C_5Me_4(SiMe_3)\lbrace 2Ti^{II} \rbrace$ (7)²⁰ (Scheme 1.1). Complex 7 was isolated by heating $\lbrace \lbrace \eta^{5} \rbrace$ $C_5Me_4(SiMe_3)\lbrace 2Ti-\eta^2-Me_3SiC=CSiMe_3\rbrace$ *in vacuo*. The paramagnetic species 6 and 7 were characterized by X-ray crystallography and revealed that the Cprings are parallel. The authors attributed the stability of **6** and **7** to the alignment and

Scheme 1.1: Examples of Well-Characterized [(η**⁵ -C5R5)2Ti(II)] Complexes**

7 6 steric bulk of the Cp-rings.

The low cost, facile synthesis, high solubility and spectroscopic properties of the benzamidinate (BTBA) ligand were attractive to Arnold and coworkers. They were able to synthesize a series of titanium (II) and titanium (III) complexes supported by (BTBA) ligands.²¹⁻ ²² The titanium (III) complex $[(BTBA)_2TiCl(THF)](8)$ was synthesized from the reduction of $[(BTBA)_2TiCl_2]$ with Na/Hg amalgam in THF (Scheme 1.2).²¹ The paramagnetic species 8 was isolated as green crystals from recrystallization in hexanes in low yield $(\leq 30\%)$. Solid-state magnetic susceptibility measurements were consistent with of a titanium (III) center. Further recrystallizations in hexanes provided paramagnetic solvent-free crystals of $[(TBA)_2Tic1]$ (9) (Scheme 1.2). Magnetic susceptibility measurements were also consistent with a titanium (III) center. The X-ray structure was in contrast to the known titanocene derivative $[Cp_2TiCl]_2$ which is dimeric. Compound **9** was found to be a good starting material for the synthesis of Ti (III) alkyls (see equation 1.2 for example). Just as in the case of the precursors **8** and **9**, the complexes were found to be paramagnetic and monomeric species.

The reduction of $[(BTBATiCl₂)]$ with Na/Hg in the presence of CO resulted in the formation of the paramagnetic Ti (III) bridging-oxo species $[(BTBA)_2Ti)_2(\mu-O)]$ (10) complex (Scheme 1.2).²² X-ray crystallography of the structure exhibits a linear Ti-O-Ti linkage. The short Ti-O bond lengths (1.821 Å) suggest partial double bond character. The magnetic susceptibility measurements were consistent with two essentially uncoupled d^1 -Ti centers.²²

The rich redox chemistry of divalent metalloporphyrin complexes led Woo and coworkers towards the synthesis of low-valent titanium porphyrin complexes.23-24 The

Me3Si

group was able to synthesize and structurally characterize the first titanium (II) porphyrin complex.23 The ligands used in this synthesis were octaethylporphyrinato (OEP) or *meso*-tetra-*p*tolyl-porphyrinato (TTP). The titanium (II) complex was synthesized from the reduction of $[(OEP)TiCl₂]$ or $[(TTP)TiCl₂]$ with LiAlH₄ in the presence of diphenylacetylene in toluene. This produced the π -alkyne titanium (II) complexes $[(OEP)Ti(\eta^2-PhC\equiv CPh)]$ (11) and $[(TTP)Ti(\eta^2-PhC\equiv CPh)]$ PhC≡CPh)] (**12**) as purple microcrystals.

 $LiAlH₄$ $[(POR)TiCl₂] + PhC \equiv CPh$ \longrightarrow $[(POR)Ti(PhC \equiv CPh)]$ **(1.3)**

> $POR = OEP (7)$ **TTP (8)**

Complexes 11 and 12 are diamagnetic and exhibit sharp ${}^{1}H$ NMR signals. The X-ray structure of **11** revealed the geometry about the Ti center was square pyramidal with the pyrrole nitrogens forming the basal plane and the alkyne triple bond at the apical site. The $C \equiv C$ stretching frequencies for the two complexes were consistent with a four-electron-donor acetylene. Woo and co-workers also synthesized a series of (TTP)Ti (II) η^2 -alkyne complexes.²³ Facile synthesis of these complexes was later developed by the reduction of $[(OEP)TiCl₂]$ or $[(TTP)TiCl₂]$ with NaBEt₃H in the presence of various alkynes.²⁴

 $[Ti(bpy)_3]$ (13), first prepared by Herzog and coworkers by the reduction of $TiCl_4$ in THF by lithium in the presence of 2-2'-bipyridyl, was isolated as a dark blue, air-and moisturesensitive compound.26 The 1H NMR spectrum showed the coordinated bipyridyl of **13** as well as the analogous $Ti(dmby)$ ₃ (14) had an upfield shift of about 1.5 ppm relative to the free \dim imine.²⁷ The upfield shift of the coordinated diimines was attributed

 $TiCl₄$ + 3 bpy **L i T H F** $[Ti(bpy)$ ³ **(1 .4)**

$$
b\,p\,y\,=\,2\,,2\,\cdot\,-b\,ip\,y\,rid\,y\,l
$$

to the presence of a considerable negative charge on the bonded ligand. Although this complex has been isolated and characterized, the exact oxidation state is not clear.

Assignments of the oxidation state of **13** and **14** have ranged from 0 to +4. Flamini and

coworkers have assigned the oxidation state of Ti for complexes **13** and **14** as +3. This suggests these complexes are best viewed as $[Ti^{3+}(byy^{-1})_3]$ although formulation of the oxidation state of the metal center has not been fully defined.28 The authors suggested complexes **13** and **14** have a singlet state far below the first excited triplet state that is not thermally accessible. An EPR signal was not given upon heating indicating a diamagnetic species. The reactivity of **13** was probed by Flamini and coworkers.^{26,27} According to the authors, the reaction of 13 with 4 equivalents of benzophenone gave the Ti(IV) complex $[Ti(bpy)(OCPh_2CPh_2O)_2]$ (15) with 2 equivalents of free bipyridine as the by-product. The author mentioned the reactivity of **13** with benzophenone was consistent of a titanium (III) complex. The authors also discovered that the addition of excess phenylisocyanate to **15** led to the catalytic formation of triphenylisocyanurate $(16)^{27}$

Scheme 1.3: Reactivity of Ti(bpy)₃

$$
16\,
$$

Recently, the synthesis of the titananorbornadiene (η^6 -arene) complex [(DMSC)Ti(1,2,4- $(Me_3Si_3C_6H_3)$] (18) (DMSC = 1,2-alternate dimethylsilyl-bridged *p-tert*-butylcalix^[4]arene) was reported.²⁸ Complex 18 was prepared from the reduction of $[(DMSC)TiCl₂]$ with Mg^{*} (activated magnesium) in the presence of excess Me₃SiC≡CH and obtained as a yellow air-and moisturesensitive diamagnetic solid in good yield. Compound 18 was characterized by ${}^{1}H$ and ${}^{13}C$ NMR and X-ray crystallography. The NMR data revealed 18 as being C_1 -symmetric in solution. Three singlets were observed for the η^6 -arene SiMe₃ group. The Bu^t groups of the calix[4]arene ligand are observed as four singlets. X-ray diffraction studies of **17** were consistent with 1,2 alternate conformation of the calix^[4]arene ligand and showed that the SiMe₃ groups are 1,2,4-

substituted around the arene ring. Although 18 can be conceived as a $Ti(II)$ - η^6 -arene the complex has significant Ti (IV) character. The complex can be more precisely described as a 7 titananorbornadiene complex. Addition of excess $\text{Bu}^t\text{C} \equiv \text{CH}$ to 18 gave $[(\text{DMSC})\text{Ti}\{\eta^6-1,3,5-\eta^6]\}$ $C_6H_3Bu_3^t$ (19). Interestingly, 19 was found to be *Cs*-symmetric in solution by ¹H and ¹³C NMR data. NMR data are consistent with the calixarene ligand of **19** being in the 1,2 alternate conformation.

The isolation and characterization of **18** provided the authors an opportunity to gain an insight of the mechanism and scope of the $[2 +2 +2]$ cycloaddition of terminal alkynes. The titananorbornadiene (η^6 -arene) complexes 18 and 19 are highly efficient catalysts for $[2 + 2 + 2]$ cycloaddition of terminal alkynes. 1,2,4-substituted benzenes were produced with excellent regioselectivity (due to the steric environment of the calix[4]arene) and in excellent yield. Compound **19** is generally less reactive toward alkynes than **18** for steric reasons. The authors found that the rate and regioselectivity of the cyclotrimerization was affected by the steric properties of the alkyne substituent. **18** and **19** were found to rarely cyclotrimerize internal alkynes. The authors noted that **18** cyclotrimerizes 65 equivalents of 2-butyne with 94% conversion. However, 93 h was required for this conversion. The potential of **18** as a (DMSC)Ti (II) synthon prompted authors to probe its reactivity with ketones and aldehydes.29 Reactions of **18** with Ar₂CO (Ar = Ph or p -C₆H₅Me) gave 2,5-dioxatitanacyclopentane derivatives $[(DMSC)Ti(OCR_2CR_2O)] (R = Ph (20) or p-C₆H₅Me (21))$ in good yields. Compounds 20 and 21 were isolated as yellow thermally stable, air-and-moisture sensitive solids.³⁰

While low-valent titanium complexes have been reported in the literature, the interest in the use of non-cyclopentadienyl ligands in LVT chemistry has grown in the past couple of decades due to the varying electronic and steric effects such ligands can impose on reactivity. This has led several groups to attempt to synthesize, isolate, and characterize LVT complexes supported by non-cyclopentadienyl complexes.³⁰⁻⁵¹ The overall goal of this research is to synthesize, characterize, and probe the reactivity of LVT complexes. The remainder of this thesis will examine the synthesis,

Ti O O Bu^t But O O Si Me Me Bu^t But R R R $[(DMSC)Ti{C_6H_3(SiMe3)3}]$ **R=SiMe3** $t-BuC \equiv CH_2$ $[(DMSC)Ti{1,3,5-C₆H₃(t-Bu)₃}]$ $xs.$ **R** \equiv H **R R R R R R + R2CO -C6H3(SiMe3)3** $-C_6H_3(SiMe_3)_3$ $(R = Ph (20) or p-C₆H₅Me (21))$ 18 $-C_6H_3(SiMe_3)$ $[(DMSC)Ti(OCR_2CR_2O)]$

Scheme 1.4: Reactivity of $[(DMSC)Ti(1,2,4-(Me₃Si)₃C₆H₃)]$

.

characterization, and reactivity of low-valent titanium bis(diimine) complexes. The synthesis was investigated in order to find a facile method of producing these complexes in good yield. Characterization was carried out to confirm the composition of the LVT bis(diimine) complexes. The reactivity with various organic substrates was probed to define the scope of the chemistry of these complexes.

CHAPTER TWO

Synthesis, structure, and reactivity of low-valent titanium bis(diimine) complexes supported by DMSC ligand

2.1 Introduction

Low-valent titanium species are of great importance in reductive coupling processes. Over the past several years, research in our group has focused on developing well-characterized low-valent titanium complexes in an effort to understand the oxidation state, structure, mechanism, and transformations mediated by low-valent titanium. The synthesis of low-valent titanium bis(diimine) complexes $[(DMSC)Ti(bpy)_2]$ (28, bpy = 2,2'-bipyridine) and $[(DMSC)Ti(dmby)_2]$ (29, dmbpy = 44'-dimethyl-2,2'-dipyridyl) was recently reported by our group.³¹ Rothwell reported the synthesis of the similar complex $[(2,6-Prⁱ₂C₆H₃O)₂Ti(bpy)₂]$ and cautiously assigned it as a $Ti(II)$ complex based on preliminary electrochemical studies.⁵¹ However, the properties and reactivity of these complexes have not been fully explored. In this chapter we would like to present our findings from our study of the properties and reactivity of **28**, **29**, and $[(DMSC)Ti(phen)_2]$ (**30**, phen = 1,10 phenanthroline).

2.2 Synthesis and characterization of [(DMSC)Ti(bpy)2], $[(DMSC)Ti(dmbpv)_2]$, and $[(DMSC)Ti(phen)_2]$

Previous work in our group has shown $[(DMSC)Ti(L_2)_2](L_2)$ = aromatic diimine) complexes 28, and 29 can be synthesized by several methods (Scheme 2.1).³⁰ The reaction of [(DMSC)TiPh2] (**27**) with at least two equivalents of the appropriate aromatic diimine has been found to be the best and most convenient method to obtain these compounds in high yield. Ozerov found that using less than two equivalents of the appropriate diimine resulted in complexes **28** and **29** with the corresponding amount of **27** remaining. This suggests that the transformation of the putative $[(DMSC)Ti(L_2)Ph_2]$ into $[(DMSC)Ti(L_2)_2]$ complex is much faster that the initial coordination of the L_2 fragment to 27^{30} Complexes 28 and 29 were found to be dark-blue, air-and moisture- sensitive solids. The compounds were found to be very soluble in aromatic hydrocarbon solvents, THF, moderately soluble in ether, and practically insoluble in pentane.

 [(DMSC)Ti(phen)2] (**30**) was similarly prepared to broaden the scope of the reactivity of the $[(DMSC)Ti(L_2)_2]$ complexes. As with the previously prepared complexes, 30 was found to be air-and moisture sensitive. Interestingly, the color of **30** was found to be olive-green instead of dark-blue as in the cases of **28** and **29**. Complexes **28**-**30** were found to be stable in the solid state but are best stored for long periods of time at low temperatures under N_2 atmosphere. Complexes 28-30 were confirmed to be C_s -symmetric in solution at ca. 22 $\rm{°C}$ by ¹H and ¹³C NMR data. Two of the six doublets integrate as two protons each and represent the methylene protons of the calix[4]arene ligand that are reflected by the mirror plane, which contains the remaining four methylene protons. The *tert*-butyl groups of the calixarene ligand are observed as two singlets each integrating as eighteen protons. In the case of **29**, the methyl protons of the dmbpy ligand were observed as two singlets (6H each). The high field shift of the *endo*-Me (located inside the calixarene cavity), compared to that of the *exo*-Me (located outside the calixarene cavity) of the bridging SiMe_2 group, gave evidence that the DMSC ligand exists in 1,2-alternate conformation (Table 2.1). The molecular structure of **28** in the solid state was determined by a single-crystal X-ray diffraction study. Selected bond angles and distances are summarized in Table 2.2.

The geometry about the Ti center is twisted away from trigonal prismatic towards

12

Scheme 2.1 Synthesis of $[(DMSC)Ti(L_2)_2]$ Complexes

| Compound | Endo-SiMe | $exo-SiMe$ |
|----------|--------------|--------------|
| | δ ppm | δ ppm |
| 28 | -0.85 | 0.29 |
| 29 | -0.78 | 0.26 |
| 30 | -0.74 | 0.34 |

Table 2.1. Selected 1 H NMR data for [(DMSC)Ti(L2)2] complexes

octahedral by about 25°, which is most likely due to the acute bite angles of the bidentate bipyridyl ligands along with the demanding steric constraints imposed by the chelating DMSC ligand. The space group for **28** is P-1. The unit cell contains one independent unit of **28** and 1.5 molecules of non-coordinated 2,2'bipyridyl to give the empirical formula $28 \cdot (C_{10}H_8N_2)_{1.5}$. Rothwell and coworkers isolated and characterized a similar complex [(2,6-

 $Prⁱ₂C₆H₃O₂Ti(bpy)₂[(31).⁵¹ The Ti-N and Ti-O bond distances of 28 are closely related to that$ of **31**. Unlike in free bipyridine, a small but systematic alternation of short and long C-C bond distances is observed within the bipyridyl ligands of **28**. The bridge C-C bonds of the bipyridyl ligands are intermediate between single and double bonds indicating disruption of aromaticity of each pyridine unit of the bipyridyl ligands. This suggests electron transfer into the LUMO of each bipyridyl ligand. Rothwell and coworkers reported similar occurrences for $[(Mo(Oprⁱ)₂(bpy)₂].⁵²]$

2.3 Reactivity of [(DMSC)Ti(L₂)₂] complexes

Photochemical and thermal substitution of an aromatic diimine ligand have been reported in the literature.^{26-27, 53-54} This left us with a desire to see if we could observe similar activity with complexes **28**-**30**. We found reactions of complexes **28-30** with two or more equivalents of Ar_2CO (Ar = C_6H_5 or Me C_6H_4) led to formation of the orange-red 1-aza-5-oxatitanacyclopentane complexes **37-42** (Scheme 2.2). Complexes **37-42** were previously synthesized in good yield by our research group from the facile reaction of one equivalent of the appropriate ketone with the titanium η^2 -ketone complexes $[(DMSC)Ti(\eta^2-OCAT_2)L_2)]$ **31-36.**⁵⁸ The transformation of complexes **31**-**36** to **37**-**42** in which C-H activation of the diimine ligand and subsequent hydride migration to a Ti-bound ketone to form an alkoxide group was

Table 2.2 Selected Bond Distances and Angles for 28

found to be photochemically assisted.⁵⁸

Complexes 37-42 were characterized by ${}^{1}H$ and ${}^{13}C$ NMR spectroscopy and microanalysis data. The X-ray structure of **40** confirmed the structure assigned for complexes **37-42** by solution NMR spectroscopy. The η^2 - ketone complexes $[(DMSC)Ti(\eta^2-OCAT_2)L_2)]$ **31**-**36** were revealed as intermediates in the reaction by 1 H NMR data. The reactions of **28**-**30** with the appropriate ketone to produce the $[(DMSC)Ti(\eta^2-OCAr_2)(L_2)]$ complexes 31-36 were found to be reversible. ¹H NMR indicated the reaction of $[(DMSC)Ti\{\eta^2-OC(\rho MeC_6H_4$)₂ ${\rm (dmbpy)}$ [(34) with one equivalent of dmbpy in C_6D_6 produced [(DMSC)Ti(dmbpy)₂] (29), 34, and the corresponding 1-aza-5- oxa-titanacyclopentene derivative $[(DMSC)Ti(\kappa^3 -$ OC(pMeC₆H₄)₂C₁₀H₅Me₂N₂}{OCH(p-MeC₆H₄)₂}] (40) in ~3:22:1 ratio after two hours whereas the reaction of $[(DMSC)Ti {\eta^2-OC(p-MeC_6H_4)_2} (bpy)]$ (32) with one equivalent of bpy in C_6D_6 produced $[(DMSC)Ti(bpy)_2]$ (28), 32, and $[(DMSC)Ti(\kappa^3-OC(p-MeC_6H_4)_2C_{10}H_7N_2)\{OCH(p-R)H_2H_3H_4H_5O(R)\}$ MeC₆H₄)₂}] (38) in ~2:2:1 ratio after two hours. The analogous reaction of [(DMSC)Ti{ η^2 - $OC(p-MeC_6H_4)_2$ (phen)] (36) and 1.4 equivalents of phen in C_6D_6 produced [(DMSC)Ti(phen)₂] (**30**) and $[(DMSC)Ti(\kappa^3-OC(p-MeC_6H_4)_2C_{12}H_7N_2)$ {OCH($p-MeC_6H_4$)₂}] (**42**) in a ~5:4 ratio in one hour. This indicates the equilibrium between $[(DMSC)Ti(L_2)]$ (28-30) and $[(DMSC)Ti\{\eta^2 - \eta^2\}]$ OC(*p*-MeC₆H₄)₂}(L₂)] (**31-36**) lies further away from **28-30** as the π-acidity of the L₂ ligand decreases. This also suggests that the reactions of **31**-**36** with aromatic diimines to form **28**-**30** occur at a faster rate than reactions of $31-36$ with Ar₂CO to produce compounds by ¹H NMR. $[(DMSC)Ti(L_2)_2 (28-30) + Ar_2CO \longrightarrow [(DMSC)Ti(\eta^2OCar_2L_2)] (31-36) + L_2$ k_2 Ar₂CO **k-1 k1** $Ar = C_6H_5$ or **(***p***-MeC6H4)2CO (eq 2.1)**

37-42

The concentration of Ar2CO also influences the rate of formation of **37**-**42** from **28**-**30**. 1 H NMR revealed that 5% of **28** was observed in solution after the reaction of **28** with two equivalents of $(p\text{-}MeC_6H_4)_2CO$ in C_6D_6 for 21 hours. However, complete consumption of 28 was observed within ~10.5 hours in the reaction of three equivalents of (*p*-MeC₆H₄)₂CO with 28 in C6D6. The rate of formation of **37**-**42** from **28**-**30** was shown to increase in the following order: bpy<dmbpy<phen. This is in contrast to the reaction of $[(DMSC)Ti\{\eta^2-OC(p MeC_6H_4$ ₂ $\{(L_2)\}$ and one equivalent of $(p-MeC_6H_4)$ ₂CO where the rate of reaction was shown to

increase in the following order: dmbpy
>bpy<phen.⁵⁸ The equilibrium between [(DMSC)Ti(L₂)₂] and [(DMSC)Ti{ η^2 -OC(p -MeC₆H₄)₂}(L₂)] complexes lies much further toward the Ti- η^2 -ketone complex when L_2 = dmbpy than when L_2 = bpy leading to greater conversion of $[(DMSC)(dmbpy)_2]$ (29) into $[(DMSC)Ti\{\kappa^3-OC(p-1)]\}$ MeC_6H_4)₂C₁₀H₅Me₂N₂)OCH(*p*-MeC₆H₄)₂}] (40) than more complete conversion of $[(DMSC)Ti(bpy)_2]$ (28) into $[(DMSC)Ti\{\kappa^3-OC(p-MeC_6H_4)_2 C_{10}H_7N_2OCH (p-MeC_6H_4)_2\}]$ (38). We found that the transformation of the $[(DMSC)Ti(L_2)_2]$ complexes **28-30** into the 1-aza-5-oxatitanacyclopentene compounds (**37-42**) is accelerated in the presence of light. For example, when three equivalents of $(p\text{-}MeC_6H_4)_2CO$ was reacted with $[(DMSC)Ti(dmby)_2]$ in the absence of light, the reaction was complete within ten hours whereas the reaction only took seven hours in ambient light.

Although the loss of a diimine from the titanium metal center is clearly involved in the reaction, the mechanistic details of the reaction of the $[(DMSC)Ti(L_2)_2]$ complexes 28-30 with two or more equivalents of ketone to yield the titanium η^2 -ketone complexes [(DMSC)Ti(η^2 - $OCAR₂(L₂)$] **31-36** are still unclear. Two scenarios can be envisioned for the reaction: i) the reaction occurs through the dissociation of a diimine prior to coordinating with the incoming ketone or ii) the incoming ketone coordinates to the metal center to give a seven-coordinate species followed by subsequent loss of a diimine.

We were surprised to find that prolonged standings of C_6D_6 or THF solutions of 28-30 in ambient light leads to formation of Ti (IV) compounds $43-45$ (Scheme 2.3). $[(DMSC)H_2]$ was consistently noticed as a minor product in this transformation. This suggests that the conversion of the $[(DMSC)Ti(L_2)_2]$ complexes 28-30 into 43-45 involves a small amount of adventitious water. To ascertain if this was the case, when a C_6D_6 solution of $[(DMSC)Ti(phen)_2]$ (30) was exposed to air (atmospheric moisture present served as the water source), the solution immediately turned orange in color and the 1 H NMR indicated that **45** was present in solution along with $[(DMSC)H_2]^{59}$ and other unidentified titanium species.

The conversion of complexes **28-30** into **43-45** appears to be photochemically assisted. When a C_6D_6 solution of $[(DMSC)Ti(phen)_2]$ (30) along with a small amount of THF was irradiated by UV light, **45** along with minor amounts of [(DMSC)H2]and unidentified titanium species in four hours. The analogous reaction in the absence of light results in no change.

$$
L_2 = 43, \text{ bpy}
$$

44, dmbpy
45, phen
Compounds 43-45 were characterized by ¹H NMR spectroscopy. The compounds were found to be C*s*-symmetric in solution. We found the calix[4]arene ligand of complexes **43**-**45** adopts an approximately cone conformation and is no longer 1,2-alternate^{29-31, 58-59} as in the case of the starting titanium bis(diimine) complexes **28**-**30**. The methylene protons of the calixarene ligand appear as four doublets. The Bu^t groups shows as three singlets integrating in a 2:1:1 ratio. The molecular structure of **45** was determined by X-ray crystallography (Figure 2.3). The structure was consistent with the ${}^{1}H$ NMR data. The titanium center adopts a distorted octahedral geometry. The X-ray structure confirms the calixarene ligand adopts a cone conformation with three of the aryloxide oxygens bound to the titanium center while the fourth is bound to the silicon atom of a SiMe₂O unit that is bound to the titanium center through the oxygen. The Ti-N and Ti-O bond distances are within the expected lengths for six-coordinate Ti(IV) complexes^{30,31} (Table 2.3).

 Complexes **28**-**30** have a potential use in organic synthesis. The hydrolysis of 1-aza-5 oxa-titanacyclopentene compounds **37**-**42** produces 2,2'-bipyridinyl-6-yl-diphenylmethanol and 1,10-phenanthroline-6-yl-diphenylmethanol compounds⁵⁹ (Scheme 2.4). Similar types of compounds have been found to be useful in various applications such as asymmetric catalysis,⁶⁰ copper-catalyzed allylic oxidation of olefins, and asymetric diethylzinc addition to aldehydes.⁶¹ The facile synthesis of $[(DMSC)Ti(L_2)_2]$ complexes 28-30 makes them attractive in synthesizing compounds similar to **46**-**51**. Complexes **28**-**30** can also be useful in understanding the role of low-valent titanium in reductive coupling of unsaturated organic substrates (i.e. ketones, aldehydes, alkynes, and imines). However, the widespread synthetic utilization of complexes **28**- **30** is hampered by the expense of the calix[4]arene ligand.

| $\overline{\text{Ti}}(1)$ -O(1) | 1.8350(15) |
|--|------------|
| $\overline{\text{Ti}}(1)$ -O(2) | 1.9404(14) |
| $\overline{\text{Ti}}(1) \text{-} \text{O}(3)$ | 1.3177(14) |
| $\overline{\text{Ti}(1)}$ -N(1) | 2.2475(18) |
| $Ti(1)-N(2)$ | 2.2540(18) |
| $Si(1)-O(4)$ | 1.6511(16) |
| $Si(1)-O(5)$ | 1.6035(16) |
| $O(1)$ -Ti(1)-O(2) | 94.81(6) |
| $O(1)$ -Ti(1)-O(3) | 105.88(6) |
| $O(2)$ -Ti(1)-O(3) | 94.08(6) |
| $O(1)$ -Ti(1)-O(5) | 97.78(7) |
| $O(1)$ -Ti(1)-N(1) | 89.82(7) |
| $O(1)$ -Ti(1)-N(2) | 162.52(7) |
| $O(2)$ -Ti(1)-N(1) | 79.54(6) |
| $O(2)$ -Ti(1)-N(2) | 80.13(6) |
| $O(2)$ -Ti(1)-O(5) | 159.64(6) |
| $O(3)$ -Ti(1)-O(5) | 97.77(7) |
| $O(3)$ -Ti(1)-N(1) | 163.57(7) |
| $O(3)$ -Ti(1)-N(2) | 91.22(7) |
| $O(5)$ -Ti(1)-N(1) | 84.54(6) |
| $O(5)$ -Ti(1)-N(2) | 83.11(7) |
| $N(1)$ -Ti(1)- $N(2)$ | 72.85(7) |
| $O(3)$ -Ti(1)-(O2) | 94.08(6) |
| $O(5)$ -Si (1) -O (4) | 111.889(8) |
| $Si(1)-O(5)-Ti(1)$ | 143.55(9) |

Table 2.3 Selected Bond Distances and Angles for 45

2.4 Experimental section

2.4.1 General details.

All experiments were performed under dry nitrogen atmosphere using standard Schlenk techniques or in a Vacuum Atmospheres, Inc. glovebox. Toluene, tetrahydrofuran, ether, and toluene were distilled from sodium benzophenone ketyl. Pentane was distilled from sodium benzophenone ketyl with addition of 1ml of tetraethyleneglycol dimethyl ether as a solubilizing agent. Benzene- d_6 was distilled from calcium hydride. All solvents were stored in the glovebox over 4A molecular sieves, that were dried in a vacuum oven at least 48 hours prior to use. Benzophenone, 1,10-phenanthroline, 4,4'-dimethylbenzophenone, 2,2'-bipyridine, and 4,4'dimethyl, 2-2'bipyridine were purchased from Aldrich Chemical Co and sublimed before use. $[(DMSC)TiCl₂](24)$, $[(DMSC)Ti(Ph)₂](27)$, $[(DMSC)Ti(bpy)₂](28)$, and $[(DMSC)Ti(dmby)₂]$ (29) were prepared as previously reported.³¹ ¹H and ¹³C NMR spectra were recorded on a Varian Gemini-200 spectrometer or a Varian VXR-400 spectrometer at ca. 22° C. ¹H and ¹³C chemical shifts were referenced to residual solvent peaks. Elemental analyses were performed by Complete Analysis Laboratories, Inc., Parsippany, NJ.

2.4.2. Preparation of Ti(diimine) complexes

 $[(DMSC)Ti(phen)Cl₂]$ (26). 1,10-phenanthroline (54.9 mg, 0.304 mmol) was added into a 10 mL solution of [(DMSC)TiCl2] (**24**) (0.250 g, 0.304 mmol) in diethyl ether. The solution was stirred at room temperature for one hour and the solvent was removed under reduced pressure to give an orange solid. The product was washed with pentane (3 x 3 mL) and dried under vacuum to give 0.266 g (87%) of product. ¹H NMR (C₆D₆) δ: 9.32 (dd, J = 4., 1.2 HZ, 2H, phen), 7.77 (d, J = 13.6 Hz, 1H, arom CH) 7.38 (d, 2H, arom CH), 7.34 (d, 2H, arom CH), 7.29 (d, J = 8.0 Hz, 2H, phen), 7.07 (d, 2H, arom CH), 6.95 (s, 2H, phen), 6.85 (dd, J = 8.0, 4.8 Hz, 2H, phen), 4.65 (d, J = 15.8 hz, 2H, calix-CH₂), 4.63 (d, 1H, calix-CH₂), 4.29 (d, J = 17.0 Hz, 2H, calix-CH₂), 3.56 (d, J = 14.0 Hz, 1H calix-CH₂), 3.38 (d, J = 15.0 Hz, 1H calix-CH₂), 1.42 (S, 18 H, t-Bu), 0.94 (s, 18H, t-Bu), 0.37 (s, 3H, *exo*-SiMe), -1.01 (s, 3H, *endo*-SiMe). 13C NMR (C₆D₆) δ: 163.5, 151.4, 148.5, 145.0, 143.3, 142.1, 141.8, 137.3, 130.7, 129.8, 129.7, 129.6, 129.3, 127.2, 127.1, 126.8, 126.7, 124.5, 40.9 (calix-CH2), 38.2 (calix-CH2), 37.5 (calix-CH2), 34.7 (*C*(CH3)3), 34.0 (*C*(CH3)3), 32.2 (C(*C*H3)3), 31.9 (C(*C*H3)3), 4.6 (*exo*-SiMe), -0.5

(*endo*-SiMe). Anal. Calcd. For C₅₈H₆₆Cl₂N₂O₄SiTi: C, 69.55; H, 6.59; N, 2.79. Found: C, 69.16; H, 6.38; N, 2.68.

 $[(DMSC)Ti(phen)_2]$ (30) . 1,10-phenanthroline $(0.617 \text{ g}, 3.42 \text{ mmol})$ was added into a 15 mL solution of [(DMSC)TiPh2] (**27**) (1.55 g, 1.71 mmol). The initially bright yellow solution immediately became dark olive-green in color. After stirring for two minutes, pentane (30 mL) was added to the solution and cooled at -15 °C for 10 minutes. The olive-green solid was isolated by filtration and washed with pentane (3 x 5 mL) and dried under reduced pressure to give 1.44 g (77%) of product. ¹H NMR (C₆D₆) δ: 10.11 (d, J = 6.0 Hz, 2H, phen), 9.25 (d, J = 6.0 Hz, 2H, phen), 7.49 (d, 2H, arom CH), 7.46 (d, 2H, arom CH) 6.95(d, 2H, arom CH), 6.84 (s, 2H, phen), 6.70 (d, J = 7.2 Hz, 2H, phen), 6.75 (d, 2H, arom CH), 6.49 (s, 2H, phen), 6.13 (d, J = 7.2 Hz, 2H, phen), 6.05 (dd, J = 7.2, 5.6 Hz, 2H, phen), 5.75 (dd, J = 7.2, 5.6 Hz, 2H, phen), 4.82 $(d, J = 16.0 \text{ Hz}, 2H, \text{ calix-CH}_2), 4.56 (d, J = 15.2 \text{ Hz}, 1H, \text{ calix-CH}_2) 4.35 (d, J = 16.0) \text{ Hz}, 2H,$ calix-CH₂) 3.92 (d, J = 13.2 Hz, 1H, calix-CH₂), 1.51 (s, 18 H, t-Bu), 0.88 (s, 18H, t-Bu),) 0.32 (s, 3H, *exo*-SiMe), -0.74 (s, 3H, *endo*-SiMe). 13C NMR (C6D6) δ: 163.6, 152.2, 150.7, 145.4, 142.8, 142.1, 140.2, 135.6, 132.1, 131.3, 130.1, 129.7, 127.6, 127.4, 127.0, 126.9, 126.5, 126.2, 125.9, 123.1, 115.8, 115.3, 41.2 (calix *C*H2), 38.7 (calix *C*H2), 38.5 (calix *C*H2O, 34.5 (*C*(CH3)3, 33.8 (*C*(CH3)3), 32.6 (C(*C*H3)3), 31.6 (C(*C*H3)3), 2.0 (*exo*-SiMe), -1.09 (*endo*-SiMe). Anal. Calcd. For $C_{70}H_{74}N_4O_4S$ iTi: C, 75.65; H, 6.71, N, 5.04. Found: C, 75.29; H, 6.57; and N, 4.95.

General Procedure for 1 H NMR Detection of Complexes 43-**45**

In a NMR tube, $[(DMSC)Ti(phen)_2] (10.0mg)$ was placed in 0.800 mL of C_6D_6 . 1 ul of THF was then added to the NMR tube. The NMR tube was placed under UV light and the transformation to 45 was monitored by ¹H NMR. Quantitative formation of 45 was formed within 36 hours along with minor amounts of $[DMSC(H_2)]$.

44 ¹H NMR (C₆D₆) δ: 8.72 (d, 2H, dmbpy), 7.40 (d, 2H, arom CH), 7.37 (d, 2H, arom CH), 7.11 (s, 2H, arom CH), 6.80 (s, 2H, arom CH), 5.96 (d, 2H, dmbpy), 5.32 (d, J = 14.0 Hz, 2H, calix-CH₂), 4.64 (d, J = 12.4 Hz, 2H, calix-CH₂), 3.72 (d, J = 14.0 Hz, 2H, calix-CH₂), 3.30 (d, J = 12.4 Hz, 2H, calix-CH₂), 1.62 (s, 6H, dbpy), 1.47 (s, 18 H, t-Bu), 1.33 (s, 9H, t-Bu), 1.04 (s, 9H, t-Bu), 0.59 (s, 6H, SiMe2) **45** ¹ H NMR (C6D6) δ: 9.12 (br d, 2H, phen), 7.45 (br m 2H, phen), 7.44 (d, 2H, arom CH), 7.40 (d, 2H, arom CH), 7.12 (s, 2H, arom CH), 6.84 (s, 2H phen), 6.77 (br s, 2H, arom CH), 6.40

(dd, 2H, phen), 5.36 (d, J = 14.0 Hz, 2H, calix-CH₂), 4.73 (d, J 12.8 Hz, 2H, calix-CH₂), 3.74 (d, $J = 14.0$ Hz, 2H, calix-CH₂), 3.35 (d, $J = 12.8$ Hz, 2H, calix-CH₂), 1.49 (s, 18 H, t-Bu), 1.32 (s, 9H, t-Bu), 1.06 (s, 9H, t-Bu), o.48 (s, 6H, SiMe₂).

2.4.3 *NMR reactions of* $[(DMSC)Ti(L_2)_2]$ *complexes* 28-30

Typical Procedure for NMR Study of the Reaction Between $[(DMSC)Ti(L₂)₂]$ **complexes 28**-**30 and 4,4'-Dimethylbenzophenone**. 0.180 mL of a 0.100 M stock solution of $(p\text{-MeC}_6\text{H}_4)_2$ CO was added to a 0.094 mL of a 0.100 M stock solution of $[(DMSC)Ti(bpy)_2]$ (28) in a screw-capped NMR tube followed by 0.518 ml of C_6D_6 . This resulted in 0.800 mL of a 0.0117 M solution of 28 and a 0.0225 M solution of $(p\text{-MeC}_6\text{H}_4)_2$ CO. The NMR tube was vigorously shaken and placed into the spectrometer at 22 °C. The reaction was monitored by recording ¹H NMR spectra immediately after inserting the sample in the spectrometer and every ten minutes thereafter. The dependence of the reaction on $(p\text{-}MeC_6\text{H}_4)_2CO$ was obtained by controlling the concentration of $(p\text{-MeC}_6H_4)_2CO$ while conducting each experiment at the same temperature using **28** as the control. The dependence of the nature of the diimine was obtained by conducting analogous reactions for **29** and **30**.

Chapter Three

Synthesis, structure, and reactivity of titanium bis(diimine) complexes supported by MBMP ligand

3.1 Introduction

The interesting reactivity of the $[(DMSC)Ti(L_2)_2]$ complexes **28-30** led us to wonder if we could synthesize and isolate similar complexes using a different ligand system. Stable and readily accessible complexes of this type could possibly be very useful in organic synthesis. We sought to find a ligand system that is i) relatively inexpensive, ii) commercially available, iii) structurally and electronically comparable to the DMSC ligand, and iv) able to form stable, monomeric titanium compounds. Our search for a ligand with the aforementioned characteristics led us to the 2,2'methylenebis(6-tert-butyl-4-methylphenaloto) (MBMP) ligand. Floriani and coworkers⁶³ reported the facile synthesis of the facile $[(MBMP)TiCl₂] (52)$ from the reaction of TiCl₄ with MBMPH₂. Floriani and Okuda⁶⁴ reported the use of 52 to synthesize stable titanium alkyl compounds. In this chapter, we present the synthesis, characterization, and reactivity of low-valent titanium bis(diimine) complexes supported by 2,2' methylene-bridged 4-methyl, 6 tert-butyl phenol ligand.

3.2 Synthesis of [(MBMP)Ti(bpy)₂], [(MBMP)Ti(dmbpy)₂], and [(MBMP)Ti(phen)₂]

Ozerov has shown the methyl groups on the titanium center of $[(DMSC)TiMe_2]$ undergo facile elimination. We were curious to see if the similar titanium complex $[(MBMP)TiMe₂]^{63-64}$ (**53**) could be utilized to prepare the desired low-valent titanium complexes. The facile synthesis and high yield of **53** were attractive to us. However, we were surprised to find **53** does not undergo reductive elimination with two equivalents of the required aromatic diimine to produce the low-valent titanium bis(diimine) complexes $[(MBMP)Ti(L_2)_2]$ (55 L₂ = bpy, 56 L₂ = dmbpy, **57** L₂ = phen) (Scheme 3.1). Instead, ¹H NMR identified complexes **58-60** as the products in the reaction.

We believed that increasing the size of the alkyl groups on the titanium center would invoke reductive elimination. To support this argument, we set out to prepare $[(MBMP)TiPh₂]$ (54) as reported in the literature.⁶³ The reaction reported by Floriani is hampered by tedious work-up and poor yields. We found that **54** could be easily isolated as a yellow solid in excellent yield ($>90\%$) by the addition of PhMgBr at low temperatures to $[(MBMP)TiCl₂] (52)$. The molecule is C_s-symmetric in solution with the plane of symmetry bisecting the TiPh₂ and containing the $-CH_2$ group as indicated by ¹H NMR data. The *tert*-butyl groups of the MBMP ligand are observed as a singlet integrating as eighteen protons. The methyl protons of the MBMP ligand are observed as a singlet (six protons). The methylene protons appear as two doublets each integrating as one proton. The compound is extremely air-and moisture-sensitive but can be stored for long periods of time at ambient temperatures under N_2 atmosphere. We found that the addition of an aromatic diimine to **54** is the most efficient method to obtaining the $[(MBMP)Ti(L_2)_2]$ complexes 55-57 in excellent yield. Compound 54 immediately turns deep dark blue upon addition of bpy or dmbpy whereas the addition of phen to **54** results in an immediate change of color to dark purple-brown. The reaction is extremely fast (complete within five minutes) and **55**-**57** can be easily isolated. The dark blue complexes $[(MBMP)Ti(bpy)_2] (55)$, $[(MBMP)Ti(dmby)_2] (56)$, and the dark brown-purple complex $[(MBMP)Ti(phen)_2]$ (57) were found to be highly air-and moisture- sensitive solids. These complexes are less soluble than the $[(DMSC)Ti(L_2)_2]$ complexes. Complexes 55 and 56 are highly soluble in THF, moderately soluble in aromatic hydrocarbon solvents and ether, and practically insoluble in pentane. Complex **57** is moderately soluble in THF, slightly soluble in aromatic solvents, and practically insoluble in ether and pentane.

Scheme 3.1: Synthesis of [(MBMP-Ti(L2)2] Complexes (**55-57**)

 The molecular structure of [(MBMP)Ti(bpy)2] (**55**) was determined by single-crystal Xray diffraction (figure 3.1). Selected bond angles and distances are summarized in Table 3.1. The geometry about the titanium center can be described as a distorted octahedral. The Ti-O and Ti-N bond distances are practically identical to those of the related complex [(2,6- $Pr^i_2C_6H_3O_2Ti(bpy)_2]^{\frac{51}{3}}(31)$ and $[(DMSC)Ti(bpy)_2]$ (28). We also noticed alternating short and long C-C bonds within each pyridine unit of the bipyridyl ligands as in the case of [(DMSC)Ti(bpy)2] (**28).**

The $[(MBMP)Ti(L_2)_2]$ complexes 55 and 56 have interesting ¹H NMR features. The ¹H NMR spectra show mostly broad peaks at ca. 22 °C. perhaps suggestive of a paramagnetic species. However, close examination of the spectra at room temperature revealed the methylene protons and the methyl groups of the MBMP ligand exhibit relatively sharp peaks. We then reasoned that the broad peaks within the ¹H NMR were due to a fluxional mechanism similar to that described in scheme 3.2. Scheme 3.2 shows an inversion of configuration by a trigonal twist via an idealized trigonal prismatic transition state. Although not as common, twist mechanisms of octahedral complexes have been reported. $53,54$

We decided to conduct variable temperature (VT) experiments on **55** and **56**. The VT-NMR experiments confirmed that **55** and **56** are fluxional in solution. Figure 3.2 shows spectra from

Figure 3.1 Molecular structure of 55

| $Ti(1)-O(1)$ | 1.882(3) |
|-----------------------|------------------------|
| $Ti(1)-O(2)$ | 1.908(3) |
| $Ti(1)-N(1)$ | 2.165(4) |
| $Ti(1)-N(2)$ | 2.216(4) |
| $Ti(1)-N(3)$ | 2.174(4) |
| $Ti(1)-N(4)$ | 2.070(4) |
| $O(1)$ -Ti(1)-O(2) | 97.27(14) |
| $O(1)$ -Ti(1)-N(2) | 95.78(14) |
| $O(2)$ -Ti(1)-N(2) | 158.86(15) |
| $O(1)$ -Ti(1)-N(3) | 165.66(16) |
| $O(2)$ -Ti(1)-N(3) | 159.86(15) |
| $N(2)$ -Ti(1)-N(3) | $\overline{76.24(14)}$ |
| $O(1)$ -Ti(1)-N(1) | 91.64(15) |
| $O(2)$ -Ti(1)-N(1) | $\overline{90.18(15)}$ |
| $N(2)$ -Ti(1)- $N(1)$ | 72.52(16) |
| $N(3)-Ti(1)-N(1)$ | 89.41(16) |
| $O(1)$ -Ti(1)-N(4) | 92.63(16) |
| $O(2)$ -Ti(1)-N(4) | 106.90(14) |
| $N(2) - Ti(1) - N(4)$ | 87.71(16) |
| $N(3)-Ti(1)-N(4)$ | 75.36(17) |
| $N(1)$ -Ti(1)- $N(4)$ | 157.57(17) |

Table 3.**1 Selected Bond Distances and Angles for 55**

Figure 3.2 Variable Temperature 1 H NMR of 55

the VT experiments for complex 55 carried out between -20° C and 10° C in toluene-d₈ (the spectra between 22 °C and 10°C were essentially identical). As the temperature is lowered, the peaks sharpen . At 10°C the tert-butyl peaks of the MBMP ligand are observed as two broad singlets. As the temperature is lowered, the peaks for the tert-butyl groups begin to sharpen*.* MBMP appear as two sharp singlets at -20 °C whereas at 10 °C the methyl groups are observed as a singlet. The bipyridyl protons could also be observed at low temperatures allowing for complex 55 to be unambiguously characterized by 1 H NMR.

At -20°C the tert-butyl groups of the MBMP ligand are observed as two singlets integrating as nine protons each. The methyl protons of the MBMP ligand also appear as two singlets (three protons each). Four signals for the MBMP ring protons are also observed. Each proton of the bipyridine ligands is inequivalent in solution. The In addition, the methyl groups of the methlyene protons appear further upfield than typical Ti complexes supported by MBMP ligand $63-64$ most likely from ring current effects of the bpy ligands. Although sharpening of peaks can be observed when solution [(MBMP)Ti(dmbpy)2] (**56**) is lowered below room temperature , the peaks in the spectra are still somewhat broad, even at -80°C. However, we were able to easily assign the peaks for the complex at -80°C.

Although further experiments must be conducted to elucidate the mechanism of the fluxional process, the VT experiments help support the twist mechanism shown in scheme 3.2. Complex **55** is *C1*-symmetric and if a fluxional process does not occur, the tert-butyl groups would be observed as two singlets. At room temperature, the tert-butyl groups are broad indicating fast conversion of the tert-butyl groups on the NMR timescale. As the temperature is lowered, the rate of conversion of the tert-butyl groups is slower on the NMR timescale and can be observed as two singlets.

The formation of complexes $55-57$ from $[(MBMP)TiPh_2]$ (54) appears to proceed in a similar fashion to that of the $[(DMSC)Ti(L_2)_2]$ complexes **28-30** (See section 2.2.1). Addition of one equivalent of bpy or dmbpy to a C_6D_6 solution of 54 resulted in the formation of either 55 or **56** with the corresponding amount of **54** remaining. When one equivalent of phen was added to **54**, the solution immediately became dark purple in color and solids formed in the bottom of the tube. However, the ${}^{1}H$ NMR of the solution revealed the presence of 54. This finding was significant because it indicates that two phen ligands coordinate to the titanium center. This, along with microanalysis data, helps confirm that **57** is formed in when two equivalents of phen

36

is added to **54**. Attempts to grow a single crystal of **57** for X-ray diffraction analysis have thus far been unsuccessful.

3.3 Reactivity of [(MBMP)Ti(L₂)₂] complexes

The $[(DMSC)Ti(L_2)_2]$ complexes 28-30 demonstrates activity towards aromatic ketones to give 1-aza-5-oxa-titanacylcopentene compounds (see scheme 2.2). The $[(MBMP)Ti(L₂)₂]$ complexes **55**-**57** exhibit similar behavior. Addition of two or more equivalents of benzophenone to a dark blue solution of [(MBMP)Ti(bpy)2] (**55**) resulted in the formation of the orange 1-aza-5-oxatitanacylcopentene derivative $[(MBMP)Ti\{\kappa^3-OC(C_6H_5)_2C_{10}H_7N_2\}\{OCH(C_6H_5)_2\}]$ (61) in good yield. Both ¹H and ¹³C NMR data for 61 are consistent with the proposed formulation. In the ¹H NMR spectrum in CD_2Cl_2 two equally intense singlets are observed for the tert-butyl groups, and two equally intense singlets for the methyl groups of the MBMP ligand. Two doublets are observed for the methylene protons of the MBMP ligand. The difference of the chemical shift between the methylene protons are much greater than in previously reported titanium compounds bearing the MBMP ligand most likely due to steric and electronic reasons. The alkoxide group hydrogen (Ti-OCH(C_6H_5)₂) of **61** is observed as a singlet at ∼δ 5.5 ppm. The ¹³C NMR resonance for the alkoxide-carbon (Ti-O-CH(C_6H_5)₂) is observed at δ 86.8 range. This is similar to the compounds previously reported by our group.58

X-ray analysis of single crystals of **61** confirmed the structure assigned by spectroscopy (Figure 3.3). The space group for **61** was assigned P 2/c. Selected bond angles and lengths are summarized in Table 3.2. The compound adopts a slightly distorded octahedral structure. The slight distortion from the idealized octahedral geometry arises from acute bite angles of the tridentate bipyridyl-alkoxide ligand [ca. 74 \degree for O(1)-Ti(1)-N(1) and 70 \degree for N(1)-Ti(1)-N(2)] in addition to the steric constraint that the bidentate MBMP ligand imposes. The Ti-N and Ti-O bond lengths are essentially identical to the analagous 1-aza-5-oxa-titanacylcopentene derivative $[(DMSC)Ti\{\kappa^3-OC(p-Me C_6H_4)_{2}C_{10}H_{7}N_{2}\}\{OCH(p-Me C_6H_4)_{2}\}]$ (40). The T-O bond distances for 61 are shorter than typical Ti-O σ -bond distances.⁵⁸ Kingston and coworkers explained the shorter Ti-O bond lengths for **40** might be reflective of a greater degree of partial Ti-O πbonding.

The $[(MBMP)Ti\{\kappa^3-OC(C_6H_5)_2C_{10}H_7N_2\}\{OCH(C_6H_5)_2\}]$ (61) and the related 1-aza-5oxa-titanacylcopentene derivatives supported by the MBMP ligand have very limited solubility. They are moderately soluble in methylene chloride, slightly soluble in chloroform, and

practically insoluble in ether, tetrahydrofuran, aromatic solvents, and pentane. However, since the reaction of $[(MBMP)Ti(bpy)_2]$ (55) and two or more equivalents of benzophenone is carried out in ether, **61** crashes out of solution upon formation allowing for facile isolation. Furthermore, any impurities can be washed away with various solvents allowing for an easy work-up to get a clean compound (based on ${}^{1}H$ NMR data). Although the reaction can be carried out at room temperature, the rate of the reaction increases upon heating up to 65°C.

The reaction of $[(MBMP)Ti(phen)_2]$ (57) with at least two equivalents of benzophenone does not readily go to the 1-aza-5-oxa-titanacylcopentene compound. Completion of the reaction took ~two weeks. Heating of the reaction while under high voltage UV lamp did not increase the rate of the reaction. The slow rate of the reaction is probably due to the high insolubility the titanium complex. The reaction between $[(MBMP)Ti(L₂)₂]$ complexes (55-57) with excess phenylisocyanate gave us interesting results. Instead of producing the expected 1-aza-5-oxatitanacyclopentene derivative, we found that complexes **(55**-**57**) readily leads to catalytic cyclotrimerization of phenylisocyanate in high yield (Scheme 3.3).^{27, 66-71} For complexes **55** and **56** the reaction with twenty equivalents of phenylisocyanate to produce the cyclotrimerized product is typically complete within twenty minutes. However, the reaction with 57 takes \sim 3 hours, due the high insolubility of the complex. The formation of $[PhNCO]_3$ (62) was established by ¹H NMR and GC/MS and was essentially identical to those values reported in the literature. Flamini reported similar reactivity with $[Ti(bpy)_3]^{27}$ (see Scheme 1.3). Initial attempts to cyclotrimerize tert-butylisocyanate have thus far failed most likely due to the steric and electronic effects the tert-butyl group of tertbutylisocyanate imposes.

In the presence of adventitious moisture or, in particular, upon exposure to air, complexes C_6D_6 solutions of 55-57 immediately turn orange. The ¹H NMR of these solutions reveals the presence the fairly air- and moisture-stable $[(MBMP)_2Ti]$ (63). Okuda⁶³ reported a similar occurrence upon deliberate hydrolysis of $[(MBMP)Ti(OiPr)_2]$. ¹H NMR showed that 63 could be formed from prolonged heating $(270^{\circ}C)$ of 55 under N₂ atmosphere. An X-ray crystal structure was obtained for the bipyridine adduct [(MBMP)2Ti(bpy)] (**64**). Complexes **55**-**57** have been shown to have similar reactivity to the $[(DMSC)Ti(L_2)_2]$ complexes **28-30**. We previously noted that DMSC 1-aza-5-oxa-titanacylcopentene compounds **37-42** have potential use in preparing produces 2,2'-bipyridinyl-6-yl-diphenylmethanol and 1,10-phenanthrolinine-6-yl-

Table 3.2 Selected Bond Distances and Angles for 61

diphenylmethanol compounds **46-51.** The cost of the calix[4]arene ligand is not favorable for widespread synthesis. We have shown that $[MBMP)Ti(L_2)_2]$ **55-57** can be used in preparing MBMP 1-aza-5-oxa-titanacylcopentene compounds. The facile preparation of complexes **55**-**57** and the low cost of the MBMP ligand are two attractive properties in the organic synthesis of produces 2,2'-bipyridinyl-6-yl-diphenylmethanol and 1,10-phenanthrolinine-6-yl-diphenylmethanol compounds. Preliminary results in our group have shown that compounds **46** and **47** can be obtained in good yield using **55**. O'Neal and coworkers have prepared similar phenanthroline compounds.⁶⁰ However, the synthesis of these compounds was limited to phenanthroline whereas we can potentially make similar bipyridine and phenanthroline compounds using **55**-**57**.

3.4 Experimental Section

3.4.1 General details

All experiments were performed under dry nitrogen atmosphere using standard Schlenk techniques or in a Vacuum Atmospheres, Inc. glovebox. Toluene, tetrahydrofuran, ether, and toluene were distilled from sodium benzophenone ketyl. Pentane was distilled from sodium benzophenone ketyl with addition of 1mL of tetraethyleneglycol dimethyl ether as a solubilizing agent. Benzene- d_6 was distilled from calcium hydride. All solvents were stored in the glovebox over 4A molecular sieves, that were dried in a vacuum oven at least 48 hours prior to use. Phenylisocyanate was purchased from Lancaster and stored over 4A molecular sieves for at least 24 hours prior to use. Benzophenone, 1,10-phenanthroline, 2,2'-bipyridine, and 4,4'dimethyl, 2-2'bipyridine were purchased from Aldrich Chemical Co and sublimed before use. 2,2' methylene-bridged 4-methyl, 6-tertbutyl phenol was purchased from Aldrich and dried under vacuum for 24 hours prior to use. $[(MBMP)TiCl₂]^{63,64}$ was prepared according to published procedures. ¹H and ¹³C NMR spectra were recorded on a Varian Gemini-200 spectrometer or a Varian VXR-400 spectrometer at ca. 22° C. ¹H and ¹³C chemical shifts were referenced to residual solvent peaks. Elemental analyses were performed by Complete Analysis Laboratories, Inc., Parsippany, NJ.

3.4.2 Preparation of Ti (diimine) complexes.

[(MBMP)TiPh2] (54) [(MBMP)TiPh2] was prepared from a modification of a known procedure.⁶³ A 15 mL ether solution of [(MBMP)TiCl₂] (52) (1.5 g 0.0033 mol) (52) was treated with PhMgBr (2.136 ml, 0.0066 mol) at -78 °C. The reaction was then allowed to warm up slowly to room temperature and stir for one hour. The solvent was removed under reduced pressure and treated with toluene. The resulted suspension was then filtered and the volatiles were removed under reduced pressure. The solid was then washed with a small amount of pentane and dried *in vacuo* to give 1.7 g (96%) of the product as a yellow solid. ¹H NMR (C_6D_6) δ: 8.23-8.25 (m, 2H Ph), 8.13-8.15 (m, 2H, Ph), 7.12-7.07 (m, 6H, Ph) 7.03 (br s, 4H, MBMParom), 3.97 (d, J = 14.4 Hz, 1H, MBMP-CH₂) 3.41 (d, J = 14.4 Hz, MBMP-CH₂), 2.12 (s, 6H, Me), 1.605 (s, 18H, t-Bu).

Typical Procedure For the Preparation of $[(MBMP)Ti(L₂)₂]$ **Complexes 55-56 To a solution** of $[(MBMP)TiPh₂]$ (3.00 g, 0.0056 mol) in 15 mL of ether was added 2-2' bipyridine (1.74 g

0.0112 mol). The reaction mixture immediately turned dark-blue and was stirred for thirty minutes. The solvent was then removed under reduced pressure and washed with pentane (3 x 10 mL) to give 3.4 g (89.5 %) of the dark-blue product.

55¹H NMR at –20 °C (toluene-d₈) δ: 10.69 (d, J = 5.2 Hz, 1H bpy), 9.20 (d, J = 5.6 Hz 1H, bpy), 8.36-8.41 (m, 2H, bpy), 7.67 (s, 1H, MBMP-arom), 7.60 (s, 1H, MBMP-arom), 7.31 (s, 1H, MBMP arom), 7.02 (s, 1H, MBMP arom), 6.68 (d, J = 8Hz, 2H, bpy), 5.89-6.00 (m, 4H, bpy), 5.85 (d, J = 12.4 Hz, 1H, MBMP-CH₂), 5.64-5.69 (m, 2H, bpy), 5.39 (t, 1H, bpy), 5.02 (t, 1H, bpy), 4.06 (d, J= 12.4 Hz, 1H, MBMP-CH2), 2.47 (s, 3H, Me), 2.40 (s, 3H, Me), 1.37 (s, 9H, t-Bu), 0.70 (s, $9H$, t-Bu). Anal. Calcd. for $C_{43}H_{46}TiO_2N_4$: C, 73.91 ; H, 6.63 ; N, 8.01 . Found: C, 73.79; H, 6.37; N, 7.92.

56¹H NMR at –80 °C (toluene-d₈) δ:10.74 (s, 1H, dmbpy), 9.22 (s, 1H, dmbpy), 8.67 (s, 1H, dmbpy), 8.52 (s, dmbpy), 7.53 (s, 1H, dmbpy), 7.28 (s, 1H, dmbpy), 7.16 (s, 1H MBMP), 7.01 (s, 1H, MBMP) 6.81 (s, 1H, MBMP), 6.60 (s, 1H, MBMP) 6.38 (s, 1H, dmbpy), 5.94 (s, CH2, MBMP), 5.59 (s, 1H, dmbpy). 5.49 (s, 1H, dmbpy), (4.027, s, CH₂), 4.02 (s, 1H, CH₂), 2.34 (s, 3H, CH3) .22 (s, 3H, CH3), 1.46 (s, 9H, t-bu), 1.39 (s, 12H, dmbpy), 0.66 (s, 9H, t-bu).Anal. Calcd. for $C_{47}H_{64}TiO_2N_4$: C, 74.78; H, 7.2, N, 7.4. Found: C, 74.62; H, 7.25; N, 7.34.

57 Anal. Calcd. for C47H46TiO2N4: C, 75.6; H, 6.2; N, 7.50. Found: C, 74.94; H, 6.46; N, 7.32. $[(MBMP)Ti\{\kappa^3-OC(C_6H_5)_2C_{10}H_7N_2\}\{OCH(C_6H_5)_2\}]$ (61). To $[(MBMP)Ti(bpy)_2]$ (0.500 g,

0.00071 mol) in ether (10 mL) was added benzophenone (0.388 g, 0.0021 mol)

The reaction was heated at 65 °C for twelve hours. The solution was then filtered and the resulting orange solid was washed with THF (3 x 5mL) and toluene (3 x 5 mL) to give 452 mg (70.5%) of the product as an orange solid.

¹H NMR (CD₂Cl₂) δ: 8.68 (d, J = 6 Hz, 2H arom CH), 8.19 (d, J = 8.4 Hz, 1H arom CH), 7.89 (t, 2H, arom CH), 7.80 (t, 2H, arom CH), 7.69 (J = 7.6 Hz, 4H arom CH), 7.58 (t, 4H arom CH), 7.35 (t, 2H, arom CH), 7.16-7.62 (m, 10 H, arom CH), 6.65 (s, 2H, MBMP), 6.48 (s, 2H, MBMP), 5.92 (d, 1H, J = 13.2 Hz, CH₂), 5.57 (s, 1H, OC*H*Ph₂), 2.73 (d, 1H, d = 12.4 Hz, CH₂), 2.19 (s, 3H, Me), 2.10 (s, 3H, Me), 1.49 (s, 9H Bu^t), 1.49 (s, 9H Bu^t).

¹³C NMR (CD₂Cl₂) δ: 170.42, 152.89, 151.32, 149.31, 148.64, 147.05, 146.60, 139.58, 138.47, 138.36, 137.29, 136.96, 135.29, 129.52, 129.42, 128.71, 128.53, 128.44, 128.20, 126.38, 126.31, 125.84, 125.71, 125.57, 125.41, 122.05, 121.89, 121.80, 119.71, 119.16, 99.54, 88.47, 86.84, 37.18, 35.54, 33.91, 31.34, 29.74, 21.70, 21.19, 20.94.

3.4.3 Cyclotrimerization of Phenylisocyanate

To $[(MBMP)Ti(bpy)_2]$ $(0.050 g)$, $(0.071 mmol)$ in toluene was added phenylisocayante 4.27 g (35.9 mmol). The reaction became dark red-brown after five minutes. The reaction was allowed to stir for one hour and filtered. The solids were washed with pentane and toluene and dried *in vacuo* to give 4.10 g of product. ¹H NMR and GC/MS data are essentially identical to the reported literature values.67-68

Chapter Four Conclusion and future research

In the course of this work, we have shown that titanium bis(diimine) complexes [(DMSC)Ti(bpy)2] (**28**), [(DMSC)Ti(dmbpy)2] (**29**), [(DMSC)Ti(phen)2] (**30**). $[(MBMP)Ti(bpy)_2]$ (55), $[(MBMP)Ti(dmby)_2]$ (56), and $[(MBMP)Ti(phen)_2]$ (57) can be easily prepared in good yield. The reactivity of the complexes was probed and we have shown that one of the diimine ligands of the titanium center can be readily displaced We have shown that the $[(DMSC)Ti(L_2)_2]$ complexes 28-30 undergoes light assisted reactions with two or more equivalents of $(p\text{-MeC}_6\text{H}_4)_2$ CO or (C_6H_5) CO to cleanly yield the corresponding 1-aza-5-oxatitanacylcopentene compound complexes. We also established that there is an equilibrium between complexes 28-30 and the Ti- η^2 -ketone complexes $[(DMSC)Ti{\{\eta^2-OC(Ar)_2\}}(31-36)$. In addition, the amount of ketone present in the reaction was shown to affect the rate of the reaction.

 The ligand about the titanium center also has an effect on the titanium bis(diimine) complexes. The $[(DMSC)Ti(L_2)_2]$ complexes have been found to be much more soluble than the[(MBMP)Ti(L₂)₂] complexes. In addition, the ¹ H NMR spectra of the [(DMSCTi(L₂)₂] complexes (**28**-**30)** display very sharp peaks and is shown to be C*s*-symmetric in solution at room temperature whereas the ¹H NMR of the $[(MBMP)Ti(L₂)₂]$ complexes 55 and 56 display broad peaks at room temperature and exhibit fluxional behavior.

The $[(MBMP)Ti(L_2)_2]$ complexes 55 and 56 are attractive in their potential uses in organic synthesis. Research in our group has indicated that complexes **55** can be a useful starting material in preparing 2,2'-bipyridinyl-6-yl-diphenylmethanol and 1,10-phenanthroline-6 yl-diphenylmethanol compounds. Complex **57** is not favorable for this reaction due to its high insolubility. Further work needs to be conducted to find favorable conditions for the use of **57.** We have also shown that complexes **55**-**57** will readily cyclotrimerize phenylisocyanate in good yield. Cyclotrimerization of phenylisocyanate using complexes **55**-**57** is advantageous over other known procedures because of the ease of synthesis of the starting materials and favorable reaction conditions. Future investigation is necessary to gain a full understanding of the mechanism of the reaction. Possible isolation of the titanium species in the reaction is also of

great interest. Reactivity with other organic substrates is needed to widen the usefulness of these complexes. In addition, preparing more titanium bis(diimine) complexes with various ligands would be useful to further examine the effect the ancillary ligand will have on the titanium center.

The oxidation states of these complexes are difficult to assign. These complexes can be viewed as Ti (II) species bound to two neutral L_2 ligands, Ti (III) species bound to one radical anionic L_2 ligand and one neutral L_2 ligand, or Ti (IV) species bound to two radical anionic L_2 ligands. It may be possible for the excited electronic state to be thermally and photochemically accessible. Theoretical data has indicated of these complexes having titanium III oxidation state. Theoretical molecular orbital calculations have been carried out on $[(MBMP)Ti(bpy)_2]$ (55) suggesting the singlet state is much higher in energy than the triplet state indicating the complex has two unpaired electrons. Evans⁷² method using ¹H NMR was conducted on complex **55.** The diamagnetic contribution⁷³ was factored in and a magnetic moment of 1.68 was found for the complex, indicative of a titanium III species. However, further EPR studies as well as magnetic susceptibility analysis are still necessary to elucidate the oxidation state of these complexes.

APPENDIX

LIST OF TABLES

Table A1. Crystal data and structure refinement for $[(DMSC)Ti(bpy)_2]$ (28)

Compound **28** Empirical formula C81H88N7O4SiTi Formula weight 1299.57 Temperature $90.0(2)$ K Wavelength 0.71073 A Crystal system, space group Triclinic, P-1
Unit cell dimensions $a = 15.1753(1)$

Volume $3438.2(5)A³$ Z, Calculated density 2 Absorption coefficient 0.199 mm^{-1} F(000) 1816 Crystal size $0.35 \times 0.15 \times 0.02$ mm Theta range for data collection 2.82 to 25.79 deg.
Limiting indices -18 -18 -18 -18 Reflections collected / unique $5846 / 6041$ [R(int) = 0.0772] Completeness to theta = 22.50 100.0 %
Absorption correction Semi-en Max. and min. transmission 0.9954 and 0.9232 Data / restraints / parameters 6041 / 7 / 568 Goodness-of-fit on F^2 1.123 Final R indices (2σ) R1 = 0.0621, wR2 = 0.1156 R indices (all data) $R1 = 0.0991$, wR2 = 0.1256 Absolute structure parameter $-0.01(4)$
Extinction coefficient $0.0017(3)$ Extinction coefficient $0.0017(3)$
Largest diff. peak and hole 0.350 and -0.253 e.A^-3 Largest diff. peak and hole

 $a = 15.1753(13)$ A alpha = 99.096(2). $b = 15.2345(13)$ beta = 113.247(2)deg. $c = 17.0936(15)$ A gamma = 101.233(2). $-18 \le h \le 18, -18 \le k \le 18, -16 \le l \le 16$ Semi-empirical from equivalents Refinement method Full-matrix least-squares on F^2

Table A3.. Crystal data and structure refinement for (45)

Compound **45** Empirical formula C81 H90 N2 O5 Si Ti Formula weight 1223.52 Temperature $150.0(2)$ K Wavelength 0.71073 A Crystal system, space group

Unit cell dimensions
 $a = 13.34700(10)$ A

Volume 7078.27(12) A^3

Z, Calculated density 4, 1.148 Mg/m³ Z, Calculated density Absorption coefficient 0.189mm⁻¹ $F(000)$ 2616 Crystal size $0.35 \times 0.30 \times 0.20$ mm Theta range for data collection 1.53 to 25.00 deg. Reflections collected / unique Completeness to theta = 22.50 100.0 %
Absorption correction Semi-en Max. and min. transmission Refinement method Full-matrix least-squares on $F²$ Data / restraints / parameters 12454 /134 / 886 Goodness-of-fit on F^2 1.037 Final R indices (2σ) R1 = 0.0474, wR2 = 0.1049 R indices (all data) $R1 = 0.0815$, $wR2 = 0.1165$ Absolute structure parameter $-0.01(4)$ Extinction coefficient $0.0017(3)$
Largest diff. peak and hole 0.385 and -0.337 e.A³ Largest diff. peak and hole

 $a = 13.34700(10)$ A alpha = 90 deg. $b = 37.2230(3)$ A beta = 110.5650 (4) deg. $c = 15.2170(2)$ A gamma = 90 deg. Limiting indices

Reflections collected / unique
 $-15 \le h \le -15$, $-44 \le k \le -44$, $-18 \le l \le -18$
 $24621 / 1254$ [R(int) = 0.0389] Semi-empirical from equivalents
0.9633 and 0.9369

Table A5. Crystal data and structure refinement for $[(MBMP)Ti(bpy)_2]$ (55)

Compound **55** Empirical formula C55 H58 N4 O2 Ti Formula weight 854.95 Temperature $90.0(2)$ K Wavelength 0.71073 A Crystal system, space group Orthorhombic, P 21 21 2
Unit cell dimensions $a = 16.9420(6)$ A alpha =

Volume 4607.0(4) A^3

Z, Calculated density 4, 1.233 Mg/m³ Z, Calculated density Absorption coefficient 0.232 mm^{-1} F(000) 1816 Crystal size $0.35 \times 0.15 \times 0.02$ mm Theta range for data collection 1.32 to 22.50 deg. Reflections collected / unique Completeness to theta = 22.50 100.0 %
Absorption correction Semi-em Data / restraints / parameters 6041 / 7 / 568 Goodness-of-fit on F^2 1.123 Final R indices (2σ) R1 = 0.0621, wR2 = 0.1156 R indices (all data) $R1 = 0.0991$, $wR2 = 0.1256$ Absolute structure parameter $-0.01(4)$ Extinction coefficient $0.0017(3)$
Largest diff. peak and hole 0.350 and -0.253 e.A^-3 Largest diff. peak and hole Max. and min. transmission

 $a = 16.9420(6)$ A alpha = 90 deg. $b = 17.5720(7)$ A beta = 90.0000(17) deg. $c = 15.4750(9)$ A gamma = 90 deg. Limiting indices

Reflections collected / unique
 $12042 / 6041$ [R(int) = 0.0772] Semi-empirical from equivalents
0.9954 and 0.9232 Refinement method Full-matrix least-squares on F^2

Table A5. Crystal data and structure refinement for [(MBMP)Ti{κ³-OC(C₆H₅)₂C₁₀H₇N₂}{OCH(C₆H₅)₂}] (61)

Compound **61** Empirical formula C65 H64 N2 O4 Ti Formula weight 985.08 Temperature 90.0(2) K
Wavelength 0.71073 A Wavelength Crystal system, space group \overline{O} Orthorhombic, P 21/c
Unit cell dimensions $a = 9.54100(10)$ A alr

Volume 5235.03(11) A^3

Z, Calculated density 4, 1.250 Mg/m^3 Z, Calculated density Absorption coefficient 0.215 mm^{-1} $F(000)$ 2088 Crystal size $0.22 \times 0.20 \times 0.15$ mm Theta range for data collection 1.55 to 25.00 deg.
Limiting indices 11<= 11, -31 < Reflections collected / unique $17990 / 9226$ [R(int) = 0.0594] Completeness to theta = 22.50 100.0 %
Absorption correction Multi-scan Absorption correction

Max. and min. transmission

Multi-scan

0.9684 and 0.9542 Max. and min. transmission Refinement method Full-matrix least-squares on $F²$ Data / restraints / parameters 9226 /189 / 679 Goodness-of-fit on F^2 1.022 Final R indices (2σ)
R indices (all data)
R indices (all data)
R indices (all data)
R indices (all data) Absolute structure parameter $-0.01(4)$ Extinction coefficient $0.0017(3)$
Largest diff. peak and hole 0.408 and -0.436 e.A⁻³ Largest diff. peak and hole

 $a = 9.54100(10)$ A alpha = 90 deg. $b = 26.3230(3)$ A beta = 95.9410(5) deg. $c = 20.9570(3)$ A gamma = 90 deg. $-11 < = h < 11, -31 < = k < 31, -24 < = k < 24$ $R1 = 0.0958$, wR2 = 0.1224

Table A9. Crystal data and structure refinement for [(MBMP)2Ti(bpy)] (**64**)

Compound **64** Empirical formula C34 H40 N O2 Ti 0.50 Formula weight 518.62 Temperature $90.0(2)$ K Wavelength 0.71073 A Crystal system, space group Monoclinic, P $2/n$
Unit cell dimensions $a = 14.2780(2)$ A

Volume 2870.05(6) A^3

Z, Calculated density 4, 1.200 Mg/m³ Z, Calculated density Absorption coefficient 0.199 mm^{-1} F(000) 1112 Crystal size $0.32 \times 0.08 \times 0.03$ mm Theta range for data collection 1.71 to 27.46 deg. Reflections collected / unique Completeness to theta = 22.50 100.0 %
Absorption correction Semi-em Max. and min. transmission Refinement method Full-matrix least-squares on $F²$ Data / restraints / parameters 6577 /0 / 347 Goodness-of-fit on F^2 1.014 Final R indices (2σ) R1 = 0.0492, wR2 = 0.11109 R indices (all data) $R1 = 0.0848$, $wR2 = 0.1245$ Absolute structure parameter $-0.01(4)$
Extinction coefficient $0.0017(3)$ Extinction coefficient $0.0017(3)$
Largest diff. peak and hole 0.486 and -0.438 e.A⁻³ Largest diff. peak and hole

 $a = 14.2780(2)$ A alpha = 90 deg. $b = 9.51900(10)$ A beta = 90.9800(5) deg. $c = 21.1200(3)$ A gamma = 90 deg. Limiting indices
 $-18 \le -18, -12 \le -12, -27 \le -12 -27$

Reflections collected / unique
 $12692 / 6577$ [R(int) = 0.0506] Semi-empirical from equivalents
0.9940 and 0.9389

References

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