## Kinetic versus Thermodynamic Metalation Enables Synthesis of Isostructural Homo- and Heterometallic Trinuclear Clusters

Sung-Min Hyun, Apoorva Upadhyay, Anuvab Das, Corey P. Burns, Siyoung Sung, Jeremy D. Beaty, Nattamai Bhuvanesh, Michael Nippe\*, David C. Powers\*

<sup>†</sup> Department of Chemistry, Texas A&M University, College Station, TX 77843, USA \*nippe@chem.tamu.edu; powers@chem.tamu.edu

**Abstract** Temperature-dependent metalation of the new hexadentate ligand (tris(5-(pyridin-2-yl)-1*H*-pyrrol-2-yl)methane; H<sub>3</sub>TPM) enables the selective synthesis of both mononuclear (*i.e.* Na(THF)<sub>4</sub>[Fe(TPM)], kinetic product) and trinuclear (*i.e.* Fe<sub>3</sub>(TPM)<sub>2</sub>, thermodynamic product) complexes. Exposure of Na(THF)<sub>4</sub>[Fe(TPM)] to FeCl<sub>2</sub> or ZnCl<sub>2</sub> triggers cluster expansion to generate homo- or heterometallic trinuclear complexes, respectively. The developed approach enables systematic variation of ion content in isostructural metal clusters via programmed assembly.

Many important multi-electron small molecule conversion processes in Biology — such as dinitrogen fixation by nitrogenases,<sup>1</sup> water oxidation by photosystem II,<sup>2</sup> interconversion of dihydrogen and protons by hydrogenases,<sup>3</sup> methane oxidation by methane monooxygenases,<sup>4</sup> and carbon dioxide reduction by Ni,Fe-CODHases,<sup>5</sup> — are accomplished by enzyme active sites that house polynuclear clusters of proximal transition metal (TM) ions.<sup>6</sup> Similarly, high-nuclearity sites at step edges and related defects have been identified as reactive sites in many heterogeneous processes.<sup>7</sup> In an effort both to better understand the intimate details of potential cooperation within polynuclear TM complexes as well as to harness the unique reactivity of high nuclearity sites for applications in catalysis, substantial effort has been directed towards the synthesis of multinuclear

TM complexes that mimic geometric or functional features of biological catalysts<sup>[8]</sup> or display new coordination geometries and electronic structures.<sup>9</sup>

Achieving predictive control over the aggregation size and geometry remains a major obstacle to the rational synthesis of polynuclear complexes.<sup>10</sup> One common approach that addresses this concern and that has been especially widely used for the preparation of trinuclear TM complexes, is the utilization of a central templating ion (such as a  $\mu_3$ -O<sup>2-</sup> ion) around which a polynuclear metal core can be stabilized (as in cores of Ru<sub>3</sub>( $\mu_3$ -O), Fe<sub>3</sub>( $\mu_3$ -O), Cr<sub>3</sub>( $\mu_3$ -O)).<sup>11</sup> Alternatively, finely-tuned, geometrically constrained, polydentate ligands can be used to exert control over critical structural aspects of polynuclear complexes. For example, the use of hexadentate ligands has proven successful in the preparation of trinuclear TM species, such as the tri-iron complexes reported by Murray and Betley (see left and right in Figure 1).<sup>9c-f</sup>



Figure 1. Comparison of recent examples of ligand-supported Fe3 complexes.

Here we describe the synthesis of a new hexadentate ligand, tris(5-(pyridin-2-yl)-1H-pyrrol-2yl)methane (H<sub>3</sub>TPM). Careful control of the metalation conditions provides access to either mononuclear (*i.e.* Na(THF)<sub>4</sub>[Fe(TPM)]) or trinuclear (*i.e.* Fe<sub>3</sub>(TPM)<sub>2</sub>, middle in Figure 1) complexes. Each of the ferrous ions in the ligand-supported trinuclear complex Fe<sub>3</sub>(TPM)<sub>2</sub> displays an unusual *cis*-divacant octahedral geometry. Further, we demonstrate that treatment of the mononuclear complex Na(THF)<sub>4</sub>[Fe(TPM)] with either FeCl<sub>2</sub> or ZnCl<sub>2</sub> results in the formation of homometallic (*i.e.* Fe<sub>3</sub>) and heterobimetallic (*i.e.* Fe<sub>2</sub>Zn) trinuclear platforms. Modular synthesis of isostructural, metal-ion substituted trinuclear complexes provides a powerful strategy to accessing systematically varied molecular clusters.

The synthesis of H<sub>3</sub>TPM was accomplished by Pd-catalyzed cross-coupling of (pyrrolyl)zinc chloride with 2-bromopyridine afforded 2-(1*H*-pyrrol-2-yl)pyridine (A), which was subsequently condensed with 1/3 equivalent of triethylorthoformate (see ESI for details). The <sup>1</sup>H NMR (Figure S1) spectrum, two-dimensional correlated spectroscopy (COSY) data (Figure S2), and <sup>13</sup>C NMR spectrum (Figure S3) are consistent with the structure of H<sub>3</sub>TPM, and high-resolution mass spectrometry confirms the molecular formula of H<sub>3</sub>TPM (HRMS (ESI<sup>+</sup>) [M+Na]<sup>+</sup> calcd. 465.1798; expt. 465.1801).

Deprotonation of H<sub>3</sub>TPM with NaHMDS at -50 °C followed by treatment with FeCl<sub>2</sub> at 23 °C affords a dark orange solution from which a red-orange crystalline solid was obtained by partial concentration and cooling to -35 °C (Figure 2a; HMDS = hexamethyldisilazide). <sup>1</sup>H NMR analysis of the resulting orange compound indicates that the three-fold symmetry of the ligand is maintained; seven paramagnetically shifted resonances are observed between -6.1 and 136.2 ppm and integrate as expected for the triply deprotonated form of the ligand (Figure S4). The <sup>57</sup>Fe Mössbauer spectrum displays a single doublet with an isomer shift ( $\delta$ ) of 0.835 mm/s and a quadrupole splitting  $|\Delta E_Q|$  of 2.171 mm/s (Figure 2b). Charge balance of three Fe<sup>2+</sup> with two triply deprotonated ligands might be expected to give rise to a trinuclear structure (Fe<sub>3</sub>(TPM)<sub>2</sub>). Consistent with such a trinuclear formulation, mass spectrometry (MALDI-TOF-MS) indicates the presence of a molecular ion with a mass of 1046.058 ([Fe<sub>3</sub>(TPM)<sub>2</sub>]<sup>+</sup> calcd. 1046.139).



**Figure 2.** Synthesis and characterization of Fe<sub>3</sub>(TPM)<sub>2</sub>. (a) Deprotonation of H<sub>3</sub>TPM with NaHMDS followed by metalation with FeCl<sub>2</sub> at 23 °C affords trinuclear complex Fe<sub>3</sub>(TPM)<sub>2</sub> in 58% yield. (b) <sup>57</sup>Fe Mössbauer spectrum of Fe<sub>3</sub>(TPM)<sub>2</sub> at 100 K which displays a single quadrupole doublet with an isomer shift ( $\delta$ ) of 0.835 mm/s and a quadrupole splitting  $|\Delta E_Q|$  of 2.171 mm/s. (c) Thermal ellipsoid plot of Fe<sub>3</sub>(TPM)<sub>2</sub> drawn at the 50% confidence interval. H atoms and co-crystallized solvent molecules have been omitted for clarity. (d) Coordination environment of the three crystallographically unique Fe ions in Fe<sub>3</sub>(TPM)<sub>2</sub>.

A single crystal of Fe<sub>3</sub>(TPM)<sub>2</sub> was obtained by cooling concentrated THF solutions at -35 °C. Refinement of X-ray diffraction data collected at 110 K results in the structure illustrated in Figure 2c. Fe<sub>3</sub>(TPM)<sub>2</sub> crystallizes in the monoclinic space group P2<sub>1</sub>/n with the whole molecule and two THF molecules residing in the crystallographically independent unit (Table S2). Each Fe centre in Fe<sub>3</sub>(TPM)<sub>2</sub> is four-coordinate and is supported by the pyridylpyrrole arms of two different ligands. The individual Fe···Fe separations vary significantly with distances of 3.041(3) Å (Fe1···Fe2), 3.203(3) Å (Fe2···Fe3), and 3.313(3) Å (Fe1···Fe3). These Fe···Fe distances are on average shorter than those observed for Murray's (µ<sub>2</sub>-H)<sub>3</sub>-Fe<sub>3</sub> complex but are substantially longer than those in Betley's Fe<sub>3</sub> complexes (2.2995(19) Å<sup>9c</sup> and 2.480(1) Å<sup>9d</sup>). The formal shortness ratio (fsr),<sup>12</sup> which is the ratio of the interatomic distance divided by the sum of the covalent radii,<sup>13</sup> for the Fe···Fe separations in Fe<sub>3</sub>(TPM)<sub>2</sub> are substantially greater than 1 (*fsr*<sub>Fe1-Fe2</sub> = 1.30, *fsr*<sub>Fe2-Fe3</sub> = 1.37,  $fsr_{Fe1-Fe3} = 1.42$ ), and thus we do not formulate any Fe–Fe bonding in complex Fe<sub>3</sub>(TPM)<sub>2</sub>. The Fe–N distances (Table 1) are comparable to those reported for previously reported four-coordinate mononuclear high-spin Fe<sup>2+</sup> (S = 2) complexes.<sup>14</sup> The metrical parameters of Fe<sub>3</sub>(TPM)<sub>2</sub> do not vary significantly with temperature from 110 K down to 10 K; see ESI for metrical parameters at 10, 30, and 50 K. Continuous shape measurement (CShM) calculations, which provide a quantitative measure of the deviation of the experimental coordination sphere from idealized geometries,<sup>15</sup> indicate that each Fe(II) center in Fe<sub>3</sub>(TPM)<sub>2</sub> adopts a nearly ideal *cis*-divacant octahedral (*i.e.* seesaw) geometry (Table 2).<sup>16</sup>

**Table 1.** Summary of geometric parameters in Fe<sub>3</sub>(TPM)<sub>2</sub>, Zn<sub>3</sub>(TPM)<sub>2</sub>, and Fe<sub>2</sub>Zn(TPM)<sub>2</sub>. Structures were acquired by X-ray diffraction at 110 K.

d (Å)	Fe <sub>3</sub> (TPM) <sub>2</sub>	Zn <sub>3</sub> (TPM) <sub>2</sub>	Fe <sub>2</sub> Zn(TPM) <sub>2</sub>
M…M	3.041(3) 3.203(3) 3.313(3)	3.5732(3) 3.6322(2) 3.7425(3)	3.3670(1) 3.4499(1) 3.5508(1)
M–N <sub>pyrrole</sub>	1.989(2)	1.951(1)	1.984(2)
M–N <sub>pyridine</sub>	2.111(2)	2.072(1)	2.078(2)

Table 2. Summary of continuous shape measurement (CShM) calculations.

Idealized	Calculated Deviation						
Geometry	Fe1	Fe2	Fe3	Zn1	Zn2	Zn3	
Square Planar	20.218	21.177	18.340	22.045	22.976	19.829	
Tetrahedral	7.428	7.416	7.690	5.528	5.523	5.897	
Cis-divacant octahedral	2.461	2.553	2.756	3.940	3.724	3.745	
Vacant trigonal bipyramid	6.896	6.631	7.655	6.254	5.981	6.948	

The trinuclear coordination mode is not unique to metalation with Fe(II). Metalation with Zn(II), accomplished by sequential treatment of H<sub>3</sub>TPM with NaHMDS and ZnCl<sub>2</sub>, affords Zn<sub>3</sub>(TPM)<sub>2</sub>. Mass spectrometry (HRMS-ESI<sup>+</sup>) was consistent with a trinuclear complex with a mass of 1073.121 (calcd.  $[M+H]^+$  1072.126) and <sup>1</sup>H NMR spectroscopy (Figure S5) indicates that

the three-fold symmetry is maintained (for <sup>13</sup>C NMR, see Figures S6).  $Zn_3(TPM)_2$  (Table S5) is isostructural to Fe<sub>3</sub>(TPM)<sub>2</sub> with longer M···M separations (3.775 Å (Zn1···Zn2, *fsr* = 1.43), 3.759 Å (Zn2···Zn3, *fsr* = 1.45), 3.684 Å (Zn1···Zn3, *fsr* = 1.50); Table 1). The Zn–N distances in Zn<sub>3</sub>(TPM)<sub>2</sub> are typical of Zn<sup>2+</sup> pyridine and pyrrolide complexes. CShM analysis indicates that each of the Zn centres in Zn<sub>3</sub>(TPM)<sub>2</sub> displays nearly ideal *cis*-divacant octahedral coordination (Table 2).

During studies of the deprotonation and metalation of H<sub>3</sub>TPM, we found that treatment of the deprotonated ligand with 1.0 equivalent of FeCl<sub>2</sub> at -80 °C (Figure 3) resulted in a <sup>1</sup>H NMR spectrum that was distinct from the spectrum obtained after metalation at 23 °C. The <sup>1</sup>H NMR spectrum obtained following low-temperature metalation features seven paramagnetically shifted resonances between -1.8 and 79.3 ppm (Figure S7). The observed peaks integrate as expected for a three-fold symmetric complex of the deprotonated ligand. We speculated that this new complex may be a monomeric complex, *i.e.* Na(THF)<sub>4</sub>[Fe(TPM)]. While we have not been able to obtain a crystalline sample of Na(THF)<sub>4</sub>[Fe(TPM)], mass spectrometry analysis reveals a major component with an *m/z* of 495.018, which is consistent with the formulation of the mononuclear complex ([Fe(TPM)]<sup>+</sup> calcd. *m/z* = 495.103).



**Figure 3.** Metalation of the deprotonated ligand with FeCl<sub>2</sub> at 23 °C affords trinuclear complex Fe<sub>3</sub>(TPM)<sub>2</sub> while metalation at -80 °C affords mononuclear complex Na(THF)<sub>4</sub>[Fe(TPM)]. Exposure of Na(THF)<sub>4</sub>[Fe(TPM)] to 0.5

equivalents of FeCl<sub>2</sub> results in the formation of  $Fe_3(TPM)_2$ . Similar cluster expansion can be accomplished by treatment of Na(THF)<sub>4</sub>[Fe(TPM)] with ZnCl<sub>2</sub>, which results in the formation of heterobimetallic cluster Fe<sub>2</sub>Zn(TPM)<sub>2</sub>.

Although once isolated, Na(THF)<sub>4</sub>[Fe(TPM)] is stable at 23 °C, treatment of a solution of Na(THF)<sub>4</sub>[Fe(TPM)] with 0.5 equivalents of FeCl<sub>2</sub> at 23 °C results in cluster expansion to generate Fe<sub>3</sub>(TPM)<sub>2</sub> in 95% yield (Figure 3). The ability to trigger the expansion of Na(THF)<sub>4</sub>[Fe(TPM)] to Fe<sub>3</sub>(TPM)<sub>2</sub> by addition of an additional 0.5 equivalents of FeCl<sub>2</sub> and the similarity of molecular structures for Fe<sub>3</sub> and Zn<sub>3</sub> complexes suggested that generation of heterobimetallic clusters from Na(THF)<sub>4</sub>[Fe(TPM)] in the presence of added Zn(II) should be feasible. Indeed, treatment of a THF-d<sub>8</sub> solution of Na(THF)<sub>4</sub>[Fe(TPM)] with ZnCl<sub>2</sub> afforded Fe<sub>2</sub>Zn(TPM)<sub>2</sub>. <sup>1</sup>H NMR analysis revealed the presence of 13 resonances (Figure S8), which would be expected from a complex in which one of the metal sites in the  $M_3(TPM)_2$  scaffold is replaced with Zn. The metal ion composition was confirmed both by ESI-MS, which provided an m/z = 1054.131 ([ZnFe<sub>2</sub>(TPM)<sub>2</sub>]<sup>+</sup> calcd. 1054.133), and by ICP-MS analysis of a HNO<sub>3</sub>-digested sample (Fe/Zn = 2.07). Crystallization from THF afforded single crystals and X-ray diffraction analysis revealed a trinuclear core that is isostructural to both  $Fe_3(TPM)_2$  and  $Zn_3(TPM)_2$  (Table S7). Each metal position is partially occupied by Fe ( $\sim 2/3$ ) and Zn ( $\sim 1/3$ ). The M–M separations (3.328, 3.407, and 3.515 Å) are intermediate between those of  $Fe_3(TPM)_2$  and  $Zn_3(TPM)_2$ . The coordination geometry at each metal is *cis*-divacant octahedral. Formation of Fe<sub>2</sub>Zn(TPM)<sub>2</sub> appears to be a kinetically controlled process and not the result of metal ion scrambling between pre-formed trinuclear clusters: <sup>1</sup>H NMR analysis of a THF-d<sub>8</sub> solution of Fe<sub>3</sub>(TPM)<sub>2</sub> and Zn<sub>3</sub>(TPM)<sub>2</sub> indicated no exchange over the course of 24 hours (Figure S9) and treatment of Fe<sub>3</sub>(TPM)<sub>2</sub> with ZnCl<sub>2</sub> results in no Zn incorporation (Figure S10).

To gain preliminary insight into the reactivity of  $Fe_3(TPM)_2$ , we have examined the redox properties of  $Fe_3(TPM)_2$  by cyclic voltammetry under an N<sub>2</sub> atmosphere. At ~ 240 mV (versus Ag/AgNO<sub>3</sub>), solutions of Fe<sub>3</sub>(TPM)<sub>2</sub> display an oxidation event that appeared to have limited reversibility. Increasing the scan rate (Figure 5) did not result in significant improvements of the reversibility of the oxidation event. Based on this observation it is likely that oxidation of Fe<sub>3</sub>(TPM)<sub>2</sub> results in a species of limited stability that undergoes chemical reaction on the time scale of the electrochemical experiment. This observation is consistent with Betley's report of the anodic oxidation of [Fe(tpe)(py)]<sup>-</sup>, which resulted in pyrrolide dissociation (tpe = tris(2-mesityl-pyrrolyl)ethane, py = pyridine).<sup>17</sup> No reversible oxidation features are observed in the CV of Zn<sub>3</sub>(TPM)<sub>2</sub> (Figures S11 and S12), which suggests significant metal-based redox chemistry in the observed oxidation event for Fe<sub>3</sub>(TPM)<sub>2</sub>.



**Figure 5.** Scan rate dependence of the initial oxidation event in cyclic voltammograms of Fe<sub>3</sub>(TPM)<sub>2</sub> under N<sub>2</sub> (0.7 mM Fe<sub>3</sub>(TPM)<sub>2</sub>, 0.1 M NBu<sub>4</sub>PF<sub>6</sub>). Potential is referenced vs Ag/AgNO<sub>3</sub>.

In summary, we report a new hexadentate ligand H<sub>3</sub>TPM. Low-temperature metalation of H<sub>3</sub>TPM with Fe(II) provides access to a mononuclear complex Na(THF)<sub>4</sub>[Fe(TPM)] as the kinetic metalation product. Metalation at ambient temperature with either Fe(II) or Zn(II) generates unusual trinuclear complexes in which each TM ion displays *cis*-divacant octahedral geometry as the thermodynamic metalation products. Exposure of Na(THF)<sub>4</sub>[Fe(TPM)] to Fe(II) or Zn(II) sources triggers the evolution of the monomeric starting material to the thermodynamically preferred trinuclear complexes. Ongoing studies are aimed at evaluating the reactivity of this new

platform of highly unsaturated trinuclear clusters towards small molecule substrates. We anticipate that the synthetic predictability that we have demonstrated for the construction of isostructural trinuclear clusters will provide access to families of structural homologs that will enable systematic evaluation of the impact on ion substitution on the reactivity of these platforms.

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