Tetranuclear Zn/4f coordination clusters as highly efficient catalysts for Friedel-Crafts alkylation

Article (Published Version)


This version is available from Sussex Research Online: http://sro.sussex.ac.uk/61500/

This document is made available in accordance with publisher policies and may differ from the published version or from the version of record. If you wish to cite this item you are advised to consult the publisher's version. Please see the URL above for details on accessing the published version.

Copyright and reuse: Sussex Research Online is a digital repository of the research output of the University.

Copyright and all moral rights to the version of the paper presented here belong to the individual author(s) and/or other copyright owners. To the extent reasonable and practicable, the material made available in SRO has been checked for eligibility before being made available.

Copies of full text items generally can be reproduced, displayed or performed and given to third parties in any format or medium for personal research or study, educational, or not-for-profit purposes without prior permission or charge, provided that the authors, title and full bibliographic details are credited, a hyperlink and/or URL is given for the original metadata page and the content is not changed in any way.
Tetranuclear Zn/4f coordination clusters as highly efficient catalysts for Friedel–Crafts alkylation†

Kieran Griffiths,a Prashant Kumar,a Geoffrey R. Akien,b Nicholas F. Chilton,c Alaa Abdul-Sada,a Graham J. Tizzard,d Simon J. Colesd and George E. Kostakis*a

A series of custom-designed, high yield, isoskeletal tetranuclear Zn/4f coordination clusters showing high efficiency as catalysts with low catalytic loadings in Friedel–Crafts alkylation are described for the first time. The possibility of altering the 4f centers in these catalysts without altering the core topology allows us to further confirm their stability via EPR and NMR, as well to gain insights into the plausible reaction mechanism, showcasing the usefulness of these bimetallic systems as catalysts.

Polynuclear coordination clusters (CCs), assembled by organic ligands, transition metal (3d) and/or lanthanide (4f) elements, represent a rapidly expanding class of materials.1–4 However, only a few and recent examples have shown the use of these materials as catalysts.5–8 Friedel–Crafts (FC) reactions9 consist of one of the most important carbon–carbon bond forming reactions in organic and medicinal chemistry. In recent years, catalysts based on zinc or lanthanide elements have been extensively employed in FC reactions and sizeable development has taken place.10,11 In particular, the indole ring system represents one of the most abundant and important heterocycles in nature exhibiting a wide range of biological activities.12 The nucleophilic addition of aldehydes or ketones to indoles, in the presence of a Lewis acid, is a facile route towards the synthesis of bisindolymethane (BIM) derivatives with only one equivalent of water generated as a side product. BIMs have recently been shown to be useful in the treatment of fibromyalgia,13 as antibacterial agents14 and even in the prevention of cancer.15 Various catalytic systems and metal salts have been reported for the synthesis of BIMs, however these methods suffer from a number of disadvantages such as expense, toxicity of reagents, high temperatures, high catalytic loadings (stoichiometric-10%) and photosensitivity (i.e. silver salts).15–18

In 3d/4f chemistry, 4f centres may be replaced by YIII or GdIII without altering the core topology,19 therefore permitting 89Y NMR20 or EPR21 for characterization of the solution species, as well as allowing study of the catalytic reaction in situ to gain mechanistic insights. Also, the possibility of altering the 3d centres while retaining the topology22,23 may provide more insights into the reaction mechanism, due to the different coordination behaviours to substrates (heteroatoms N, O or S), anticipated to be seen when Co, Ni, Cu, or Zn is used. We recently committed to studying the catalytic properties of tetranuclear 3d/4f CCs possessing a rigid defective dicubane topology assembled solely by the Schiff base organic ligand (E)-2-(2-hydroxy-3-methoxybenzylidene-amino)phenol) H1L (Scheme S1, ESI†). We showed that a series of isoskeletal24 M4III6LnII1 (M = Co or Ni, Ln = Y, Nd, Eu, Gd, Tb, and Dy) CCs effectively catalyse a domino reaction at room temperature.6,25 These heterometallic species remain intact in organic solvents and this precise topology brings the 3d and the 4f centres very close (approximately 3.3 Å), allowing both metal centres to coordinate to the substrates and promote the coupling reaction in high yield, highlighting the catalytic utility of this tetranuclear bimetallic motif.

Motivated by our previous results, and having in mind that Ln(OTf)3-26 and Zn(ClO4)2/Schiff base27 compounds have been employed as catalysts in the reaction of indole derivatives with aldehydes and ketones with a loading of 10 mol%, we decided to synthesize and characterize a series of isoskeletal Zn2III6Ln2II CCs and study, for the first time, their application in FC reactions.

The combination under aerobic conditions of H2L with freshly prepared Ln(NO3)3·xH2O and Zn(NO3)2·6H2O in the presence of Et3N in EtOH afforded, in very good yields, precipitates subsequently crystallized by vapour diffusion of Et3O in N,N′-DMF solutions, affording isoskeletal air-stable tetranuclear defective dicubane compounds with the general formula [Zn2III6LnII3(NO3)2(DMF)2]2 where Ln is Y (1Y), Sm (1Sm), Eu (1Eu) Gd (1Gd), Dy (1Dy), Tb (1Tb) and Yb (1Yb) (Fig. 1). These compounds were characterized in full by X-ray crystallography (Tables S1 and S2, ESI†), IR, ESI-MS, TGA and elemental analysis (see the ESI†).
The Zn$^{14}$ ion adopts a slightly distorted octahedral geometry with an O$_2$N donor set, while the Ln ion has a very distorted square antiprismonic geometry, by virtue of chelating nitrate anions, with an O$_2$N donor set. Five out of six (Zn) and six out of eight (Ln) coordination sites are occupied by donors from the organic ligands. The remaining coordination sites are occupied by one DMF molecule (Zn) and one nitrate anion (Ln, chelated mode). For 1Y, the Y...Zn distances are 3.34882(7) and 3.5272(8) Å and the Zn...Zn distance is 3.2949(8) Å; the other 1Ln species are similar.

The identity of 1Ln in solution was confirmed by electrospray ionization mass spectrometry (ESI-MS) studies; in all cases, we observed two peaks in the MS (positive-ion mode) at $m/z$ which corresponds perfectly to the monocationic [Zn$_2$II Ln$_3$III $\text{L}_n$(NO$_3$)$_2$]$^+$ and the dicationic [Zn$_2$II Ln$_3$III $\text{L}_n$]$^{2+}$ fragments, respectively (see Fig. S1, ESI†).

For further confirmation of the solution stability of 1Ln, Q-band EPR studies of 1Gd in both solid and solution (80% DMF and 20% Et$_2$O) phases were performed, Fig. 2. Simulations with PHIP$^*$ confirm that the spectra owe to $S = 7/2$ Gd$^{3+}$ ions with rhombic zero-field splitting (ZFS) (see Fig. S1, ESI†). The highly sensitive, finger-print-like ZFS of the $S = 7/2$ state directly indicates that the coordination environment of the Gd$^{3+}$ ion remains unchanged in solution.

For benchmarking studies, indoles and benzaldehyde were selected as reactants using 1Dy as the catalyst at 10% loading, and the reaction parameters were subsequently optimized. The first set of reactions was performed in order to identify the ideal solvent system; after screening several solvents (Table S3, ESI†) we identified that the ethanol/water (2/1) solvent system (Table 1, entry 2) produced only slightly lower yields, while zinc triflate (Table 1, entry 5) showed poor catalytic performance. Compound 1Dy afforded almost quantitative yields of the desired product after 2 h at 10 mol% loading. By use of 1Sm, 1Eu, 1Tb, 1Gd, and 1Yb comparable performance to 1Dy was obtained, however the use of 1Y at 2.5 mol% loading afforded the desired material in quantitative yield (Table 1, entry 10). Promisingly, the catalyst loading for 1Y could be decreased to 1 mol% with only a slight decrease in the yields (Table 1, entry 11), being far lower than other reported systems. Importantly, reducing the catalyst loading for yttrium triflate to 2.5 mol% (Table 1, entry 4) resulted in comparable yields.

For benchmarking studies, indoles and benzaldehyde were selected as reactants using 1Dy as the catalyst at 10% loading, and the reaction parameters were subsequently optimized. The first set of reactions was performed in order to identify the ideal solvent system; after screening several solvents (Table S3, ESI†) we identified that the ethanol/water (2/1) solvent system (Table 1, entry 2) resulted in the optimum solvent system, we conducted reactions without the 1Dy catalyst (Table 1, entry 1) and no product was obtained. Next, due to the heteronuclear (Zn/Ln) character of our molecules, lanthanide and zinc triflates were utilized at 10 mol% loading (Table 1, entries 2–5), in order to identify their influence on the bisindolylalkane product. Yttrium triflate (Table 1, entry 3) was found to give quantitative yields and dysprosium triflate (Table 1, entry 2) produced only slightly lower yields, while zinc triflate (Table 1, entry 5) showed poor catalytic performance. Compound 1Dy afforded almost quantitative yields of the desired product after 2 h at 10 mol% loading. By use of 1Sm, 1Eu, 1Tb, 1Gd, and 1Yb comparable performance to 1Dy was obtained, however the use of 1Y at 2.5 mol% loading afforded the desired material in quantitative yield (Table 1, entry 10). Promisingly, the catalyst loading for 1Y could be decreased to 1 mol% with only a slight decrease in the yields (Table 1, entry 11), being far lower than other reported systems. Importantly, reducing the catalyst loading for yttrium triflate to 2.5 mol% (Table 1, entry 4) resulted in comparable yields.

Table 1 Comparison of catalytic activity for Ln(OTf)$_3$ and 1–3–Ln catalysts

<table>
<thead>
<tr>
<th>Entry</th>
<th>Catalyst</th>
<th>Loading$^a$/mol%</th>
<th>Time/h</th>
<th>Yield$^c$/%</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>None</td>
<td>N/A</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>Dy(OTf)$_3$</td>
<td>10</td>
<td>12</td>
<td>95</td>
</tr>
<tr>
<td>3</td>
<td>Y(OTf)$_3$</td>
<td>10</td>
<td>12</td>
<td>Quantitative</td>
</tr>
<tr>
<td>4</td>
<td>Y(OTf)$_3$</td>
<td>2.5</td>
<td>12</td>
<td>55</td>
</tr>
<tr>
<td>5</td>
<td>Zn(OTf)$_3$</td>
<td>10</td>
<td>12</td>
<td>58</td>
</tr>
<tr>
<td>6</td>
<td>1Dy</td>
<td>10</td>
<td>2</td>
<td>Quantitative</td>
</tr>
<tr>
<td>7</td>
<td>1Dy</td>
<td>5</td>
<td>2</td>
<td>96</td>
</tr>
<tr>
<td>8</td>
<td>1Dy</td>
<td>2.5</td>
<td>12</td>
<td>92</td>
</tr>
<tr>
<td>9</td>
<td>1Dy</td>
<td>1</td>
<td>12</td>
<td>87</td>
</tr>
<tr>
<td>10</td>
<td>1Y</td>
<td>2.5</td>
<td>12</td>
<td>Quantitative</td>
</tr>
<tr>
<td>11</td>
<td>1Y</td>
<td>1</td>
<td>12</td>
<td>96</td>
</tr>
<tr>
<td>12</td>
<td>1Sm</td>
<td>2.5</td>
<td>12</td>
<td>87</td>
</tr>
<tr>
<td>13</td>
<td>1Eu</td>
<td>2.5</td>
<td>12</td>
<td>53</td>
</tr>
<tr>
<td>14</td>
<td>1Tb</td>
<td>2.5</td>
<td>12</td>
<td>80</td>
</tr>
<tr>
<td>15</td>
<td>1Gd</td>
<td>2.5</td>
<td>12</td>
<td>75</td>
</tr>
<tr>
<td>16</td>
<td>1Yb</td>
<td>2.5</td>
<td>12</td>
<td>94</td>
</tr>
</tbody>
</table>

$^a$ Reaction conditions: indole, 1 mmol; benzaldehyde, 0.5 mmol; catalyst; 10 mL EtOH/H$_2$O (2:1); room temperature. $^b$ Catalyst loading calculated per equivalent of Ln$^{3+}$. $^c$ Determined by $^1$H NMR spectroscopy.
in a severe decrease in yield, compared to the ability of 1Dy and 1Y to maintain high yields at only a few mol%.

We then explored the scope of the reaction by employing a variety of aldehydes and substituted indoles (Table 2). The reaction proceeds smoothly with very good to excellent yields. Products 7a, 8c and 8f were characterized via single crystal X-ray diffraction (Fig. S20, ESI†). The next step was to involve ketones in place of aldehydes. Long reaction times even with a slight increase of the temperature (50 °C) did not yield any product with both aliphatic (acetone, cyclohexanone) and aromatic ketones (acetophenone) and no unexpected side products were observed. This opposite behaviour recalls the selective reduction of ketones in the presence of aldehydes (Luche reaction).29

Table 2 1Y catalysed FC alkylation of indoles and aldehydes

<table>
<thead>
<tr>
<th>R1</th>
<th>R2</th>
<th>Product</th>
</tr>
</thead>
<tbody>
<tr>
<td>H</td>
<td>H</td>
<td>7a, 100%</td>
</tr>
<tr>
<td>O,</td>
<td></td>
<td>7b, 95%</td>
</tr>
<tr>
<td>7c</td>
<td></td>
<td>7c, 97%</td>
</tr>
<tr>
<td>7d</td>
<td>1%</td>
<td>7d, 96%</td>
</tr>
<tr>
<td>7e</td>
<td>O,</td>
<td>7e, 89%</td>
</tr>
<tr>
<td>7f</td>
<td>O,</td>
<td>7f, 79%</td>
</tr>
<tr>
<td>7g</td>
<td>7%</td>
<td>7g, 72%</td>
</tr>
<tr>
<td>7h</td>
<td>7%</td>
<td>7h, 65%</td>
</tr>
</tbody>
</table>

Reactions conditions: aldehyde, 0.5 mmol; substituted indole, 1 mmol; EtOH/H2O (2:1) 10 mL; room temperature; 12 h. 2.5% loading of 1Y.

In this reaction, aldehydes bond to Ln centres via hemiketal form, whilst ketone remains unprotected, and is thus reduced by NaBH4. Efforts to monitor the formation of the hemiketal form by NMR were not successful. To gain information on the reaction mechanism we performed UV-Vis binding studies of 2-naphthaldehyde with Zn(OTf)2, Dy(OTf)3, and 1Dy (Fig. S21, ESI†). 1Dy is found to have almost double the rate of 2-naphthaldehyde quenching compared to Dy(OTf)3 and Zn(OTf)2 is significantly slower than both. This observation indicates that 2-naphthaldehyde prefers coordination to Dy over Zn.

To gain further information on the plausible mechanism and to identify the limitations of the present catalysts, we decided to perform the following set of reactions. First, we employed an aliphatic aldehyde (cyclohexanecarbaldehyde) instead of an aromatic aldehyde in reaction with indoles, which gave very low yields of the expected product 9 (Scheme S1, ESI†) after 72 hours. Second, we examined substituted indoles such as 2-methyl-indole, 2-(trifluoromethyl)-indole, 3-methyl indole, indole-3-acetic acid and N-methyl indole in reaction with benzaldehyde. The use of 2-methyl-indole gave the expected product 10 (Scheme S1, ESI†) in quantitative yield in only 2 hours, indicating that substitution in position 2 promotes the reaction. Compound 10 was characterized via single crystal X-ray diffraction (Fig. S22, ESI†). However, the use of the electron withdrawing group –CF3 in place of –CH3 [reaction with 2-(trifluoromethyl)-indole] gave a yield of only 16%, product 11, indicating a substantial influence on the catalytic activity. The use of indole-3-acetic acid did not yield any product. A logical explanation for deterioration of catalytic activity is that the carboxylic group of indole-3-acetic acid is in competition with benzaldehyde to coordinate to the Ln centre, leading to a poisoning of the catalyst, however the use of 3-methyl-indole did not result in the formation of BIM. The latter indicates that if the most active site (C-3) of the indole group is blocked the reaction does not proceed. Finally, the reaction of N-methyl-indole with benzaldehyde did not yield any product, showcasing that coordination of the nitrogen atom is crucial.

Despite the diamagnetic nature of 1Y, characterizing it in the solution state by NMR proved to be difficult due to its asymmetry, dynamic behaviour, and its relatively low solubility. However, the peak areas for the imine, aromatic, and methoxy protons were all consistent with the structure as determined by X-ray crystallography, Fig. 1. Gradually warming to 75 °C in 10 °C steps caused gradual broadening of the peaks, which then recoalesced upon cooling back to room temperature, without any apparent decomposition of the complex. Using multiple solvent suppression, it was possible to monitor the smooth conversion of benzaldehyde and indole to the product in protonated 2:1 EtOH/H2O. With Y(OTf)3, the concentration of intermediates appeared to be very low, but with prolonged acquisition it was possible to identify a minor species using 13N-HMBC. Its chemical shift of 113 ppm is distinctly different from indole (130 ppm) and the product (125 ppm), but the proton chemical shifts of 7.33 (127.9 ppm in 13C-HSQC) and 4.99 ppm (too close to the solvent peak to identify 13C-correlations) suggest that it may be the benzaldehyde-indole hemiaminal. In the absence
of a catalyst, the product forms very slowly (ca. 10% in 1 month), but under the same spectroscopic conditions it was not possible to detect the same correlations. This raises the prospect that it may well be a key intermediate, but more favourable conditions may be required to unambiguously identify it.

Given knowledge of the structure of the catalyst and the data provided above, we propose the following mechanism for the FC reaction (Scheme 1). The first step of the reaction involves the coordination of the N\textsubscript{indole} atom to the \textit{Zn}\textsuperscript{2+} and the O\textsubscript{carboxyl} atom of the aldehyde to the Ln part of the catalyst (Step 1). In the presence of H\textsubscript{2}O the aldehyde may coordinate to the Ln centre \textit{via} its hydrated form. Then, deprotonation of the coordinated indole leads to the formation of a negative centre at C-3 of the indole moiety (Step 2). The two organic moieties are very close [Y - Zn distances are 3.34882(7) and 3.5272(8) À] to favour the formation of benzaldehyde-indole hemiaminal, as suggested by the NMR studies (Steps 3 and 4) followed by alkylation of one more indole moiety (Step 5). Finally, the catalytic cycle ends with a proton exchange with an additional indole moiety (Step 6) and release of the bis-adduct product and water and reform of the active catalyst species (Step 7).

Overall, the results presented herein confirm for the first time that these tetranuclear CCs may indeed be useful as FC catalysts, with low catalyst loadings. Our philosophy to alter the 4f elements in these solution-stable bimetallic 3d/4f species allow us to confirm their stability \textit{via} EPR and gain useful insights into the plausible reaction mechanism \textit{via} NMR. Moreover, the possibility of tuning the organic periphery of these catalysts,\textsuperscript{16,17} achieving immobilization, showcases that the present catalysts are highly desirable and appropriate vehicles to open new avenues in the fields of 3d/4f chemistry and catalysis. Further work is necessary to expand the scope of the present Zn/Ln CCs to other FC alkylation reactions, and this will be the focus of our future activities.

We thank the EPSRC (UK) for funding (grant number EP/M023834/1), the University of Sussex, for offering a PhD position to K. G., the EPSRC UK National Crystallography Service at the University of Southampton\textsuperscript{10} for the collection of the crystallographic data for compounds 7\textit{a}, 8\textit{c}, 8\textit{f} and 10, and the EPSRC UK National Electron Paramagnetic Resonance Service at the University of Manchester.

Notes and references

\begin{itemize}
\itemATALytic protocol. To an EtOH/H\textsubscript{2}O (2:1 in 10 mL) mixed solution, aldehyde (0.5 mmol) and indole (1 mmol) were added followed by the catalyst (2.5% with respect to aldehyde). The resultant solution was stirred for 12 h, during which time the product had precipitated. The cloudy solution was filtered and the precipitate was washed with hexane (3 × 10 mL) and water (3 × 10 mL). For products which did not precipitate, the clear solution was concentrated under reduced pressure and extracted in ethyl acetate (30 mL) from water (20 mL). The ethyl acetate layer was dried with MgSO\textsubscript{4} and concentrated under reduced pressure. The resulting oil was washed with hexane (60 mL), resulting in a powdered solid which was further washed with water (3 × 10 mL).


\end{itemize}