This is an Accepted Manuscript, which has been through the Royal Society of Chemistry peer review process and has been accepted for publication.

Accepted Manuscripts are published online shortly after acceptance, before technical editing, formatting and proof reading. Using this free service, authors can make their results available to the community, in citable form, before we publish the edited article. We will replace this Accepted Manuscript with the edited and formatted Advance Article as soon as it is available.

You can find more information about Accepted Manuscripts in the Information for Authors.

Please note that technical editing may introduce minor changes to the text and/or graphics, which may alter content. The journal’s standard Terms & Conditions and the Ethical guidelines still apply. In no event shall the Royal Society of Chemistry be held responsible for any errors or omissions in this Accepted Manuscript or any consequences arising from the use of any information it contains.
Tungsten imido catalysts for selective ethylene dimerisation

Christopher M. R. Wright, Zoë R. Turner, Jean-Charles Buffet and Dermot O’Hare*

A tungsten imido complex W(NDipp)MeCl (Dipp = 2,6-Pr₂C₆H₄) is active for the selective dimerisation of ethylene to yield 1-butene under mild conditions. Immobilisation and activation of W(NDipp)Cl₄(THF) on layered double hydroxide, silica or polymethylaluminoxane yields active solid state catalysts for the selective dimerisation of ethylene. The polymethylaluminoxane-based catalyst displays a turnover frequency (4.0 molₖg⁻¹mol⁻¹h⁻¹) almost 7 times that of the homogeneous catalyst.

Cheap feedstocks (methane, ethane) and energy derived from shale gas exploration has led to a $135 billion investment in the US petrochemical industry. Processes that upgrade short chain α-olefins produced from such resources, especially with a high level of control, are therefore of great value. The last decade has seen the development of highly selective ethylene oligomerisation catalysts of which dimerisation and trimerisation are most prevalent. Currently, the most selective and productive systems are chromium-based, however, such high levels of selectivity are still difficult to achieve or predict. Perhaps the most famous industrial method is Axens’s AlphaButol process based on a titanium catalyst which is high yielding (~93% conversion in 15 minutes at 2 MPa ethylene pressure) and highly selective for C₂ olefins (>99%). Tungsten-based systems have also been shown to be efficient olefin dimerisation catalysts although to a much lesser extent. The most prominent of these involve N-donor ligands and are formed either in situ from WCl₆ and anilines or by activating tungsten imido complexes with aluminium alkyls; >99% selectivity towards dimer formation has been demonstrated.

Making molecular catalytic systems heterogeneous has many industrial advantages such as ease of product separation and catalyst recycling. Well-defined tungsten catalysts for alkene and alkane metathesis have been primarily based on silica supports, using the concept of surface organometallic chemistry (SOMC) to immobilise the complexes. Basset and co-workers have reported that the highly reactive tungsten methyl bonds in WMe₅ can be controlled when supported on partially dehydroxylated silica to produce a catalyst that is active for the metathesis of linear alkanes at elevated temperatures. Examples of supported oligomerisation catalysts in the literature are rare, however, Wass and co-workers have described how a PNP pincer chromium complex formed from CrCl₃ and (C₆H₄(OMe))₃PN(Me)(C₆H₄(OMe))₂, which is active for ethylene trimerisation with >99% selectivity, could be successfully supported on MAO-modified silica with no apparent loss of activity or selectivity. Layered double hydroxides (LDHs), represented by the formula [M⁺₃−₅xM₂⁺x(OH)₂]y(OMe)₃₋ₓ/₂yH₂O, have been shown to act as effective catalyst supports. Recent work by O’Hare and co-workers has focused on supporting olefin polymerisation complexes on LDHs. Here we report a well-defined tungsten imido complex which is active for the selective dimerisation of ethylene to 1-butene, both in solution and when supported on layered double hydroxides, silica or polymethylaluminoxane.

The complex W(NDipp)Me₂Cl (Dipp = 2,6-Pr₂C₆H₄) can be prepared from the reaction of W(NDipp)Cl₄(THF) (L = THF or Et₂O) and trimethylaluminoxane (TMA) (Scheme 1). This reactivity is analogous to that reported by Dyer and co-workers to form the bis-imido complex W(NDipp)AlMe₂(μ-Cl)(NDipp)Me₂, however, in the case of
W(NDipp)MeCl no aluminium coordination is observed. Excess TMA does not result in further methylation at ambient temperature, and complex formation also occurs when the starting material is reacted with an excess of methylaluminoxane (MAO) or dimethylaluminium chloride (DMAC).

Analysis of the molecular structure of W(NDipp)MeCl (Fig. 1), shows that it adopts a trigonal bipyramidal arrangement with the methyl groups sitting in the equatorial plane. This orientation was also proposed by Pedersen et al. for the analogous complex W(NPh)MeCl, which was not structurally characterised.\(^1\) The W1-N1-C4 angle in W(NDipp)MeCl shows a deviation away from linearity (171.0(5)° vs 178.4(3)° in W(NDipp)Cl\(_2\)(THF)) due to steric repulsion between the methyl and isopropyl substituents. This distortion is accompanied by an increase in the tungsten nitrogen bond length (1.750(8) Å vs 1.721(4) Å).

\(^1\)H NMR spectroscopy displays a singlet resonance for the three methyl groups at 1.41 ppm with tungsten satellites (\(\delta_{WH} = 7.03\) Hz). Density functional theory (DFT) calculations at the BP86 level of theory were carried out on W(NDipp)MeCl and the HOMO shows a \(\pi\)-bonding interaction between the W1 \(d_y\) and the imido N1 \(p_y\) orbitals (Fig. S4).

A solution of W(NDipp)MeCl in benzene is active for the selective dimerisation of ethylene to 1-butene in the presence of MAO (Scheme 2). Reactions carried out at 100 °C in \(\text{C}_6\text{D}_6\) with an internal standard show that lower W:MAO ratios result in greater activity and selectivity for 1-butene; isomerisation to 2-butenes is almost completely eliminated for the 1:2 ratio (Fig. 2 and S6), with a total ethylene conversion of 88% and a 95% selectivity for 1-butene after 44.5 hours.

![Image](image-url)

**Scheme 2.** Dimerisation of ethylene to 1-butene catalysed by W(NDipp)MeCl and MAO.

![Image](image-url)

**Fig. 1.** Molecular structure of W(NDipp)MeCl. Thermal ellipsoids are drawn at 50% probability. Hydrogen atoms omitted for clarity.

![Image](image-url)

**Fig. 2.** Turnover (TON) as a function of time for varying W(NDipp)MeCl:MAO ratios at 100 °C in \(\text{C}_6\text{D}_6\). [W(NDipp)MeCl]= 4.55 μmol.

\(^1\)H NMR spectroscopy shows that as the co-catalyst ratio is increased (from 1:2 to 1:40), the orange solution (1:2, 1:5 and 1:10) darkens to a brown colour (1:20) and then a dark brown (1:40) which corresponds to an intractable mixture of products which are unreactive towards ethylene (Fig. S12). A closer study of the product distribution for the 1:5 ratio (Fig. S9) shows that minor complex decomposition results in increased isomerisation to yield 2-butenes. In the presence of ethylene and 2-butenes the complex becomes active for the metathesis reaction to form propylene, a common transformation for complexes of this motif,\(^3\) indicating that an alkylidene may form in situ.

Addition of other aluminium alkyls has a marked effect on the reactivity. In the presence of DMAC (W:DMAC = 1:2) resonances for the starting complex are unaffected yielding an active catalyst for dimerisation, however, addition of TMA (W:TMA = 1:2) and subsequent heating at 100 °C results in a dark brown solution with multiple resonances exhibiting tungsten satellites in the \(^1\)H NMR spectrum, which is unreactive towards ethylene (Fig. S14). Reaction of W(NDipp)MeCl with MeLi results in a colour change from orange to yellow followed by rapid decomposition (no observable resonances in the \(^1\)H NMR spectrum) as described for W(NPh)MeCl.\(^3\) In both cases we propose that an unstable tetramethyl species is formed which decomposes via a transient methylidene, explaining the observed metathesis chemistry from ethylene to propylene which is observed at higher temperatures.

It has been suggested that the high level of selectivity seen in tungsten systems may be a result of a metallocyclic mechanism, as seen in chromium systems.\(^7\) Tobishop reported a DFT study on the role Lewis acids play in \(\alpha\)-olefin oligomerisation catalysed by mono(imido) tungsten complexes via a metallocyclic mechanism.\(^3\) The complex W(NPh)R\(_2\)Cl\(_2\) (R = olefin), proposed as the active species formed in situ, was used in the calculations and the Lewis acid (AlMe\(_3\)Cl) was shown to decrease the activation energy irrespective of the coordination mode. Our results indicate that the in situ
precatalyst may actually be a trimethyl complex W(NPh)Me₃Cl which then coordinates ethylene (Scheme S4). Mixed C₂H₄/C₃D₈ labelling studies on bis(imido) species and on an in situ tungsten initiator suggest a Cossee-type coordination-insertion mechanism may be involved, due to significant scrambling of the resulting oligomers.⁷ ³⁷ The hydride species involved can lead to hydrogenation and isomerisation of the starting materials and products. Whilst we do see ethane and butane for all W:Al ratios, these along with isomerisation to 2-butenes of the product are significantly reduced with lower cocatalyst ratios.

Utilising the SOMC concept,¹⁰ W(NDipp)Cl₃(L) (L = THF, Et₂O) can be supported on the surface hydroxyl groups of layered double hydroxides (LDHs) and silica, or on MAO-modified LDHs or polymethylaluminoxane (Scheme 3). Mixing of a green solution of complex in toluene (5% by weight) with a suspension of the solid support in toluene at ambient temperature for one hour resulted in a colourless solution and coloured support indicating immobilisation of the complex had occurred. Spectroscopic evidence indicates that methylation of the complex occurs in situ when grafted on an MAO-modified or polymethylaluminoxane support. A solution of H NMR spectrum of the polymethylaluminoxane catalyst in C₆D₆O revealed a small quantity of leached complex with resonances corresponding to W(NDipp)Me₂Cl (Fig. S45).

¹³C and ²⁷Al solid state CPMAS NMR spectroscopic studies of the supported catalysts confirmed the presence of the disoproplylphenyl groups on the surface (Figs. S17-31). The ³⁷Al NMR spectra of the LDH-supported species contain a small resonance for the surface aluminium species at ~77 ppm, relative to the resonance at ~8 ppm representing the bulk aluminium environment of the LDH.

Oligomerisation studies in C₆D₆ with MAO as a co-catalyst revealed that W(NDipp)Cl₃(L) (L = THF, Et₂O) supported on silica, Mg₃Al₂O₄ and Mg₃Al₂O₃ LDHs to be active heterogeneous catalysts for ethylene dimerisation, whereas the Mg₃Al₂O₄ only formed a small amount of polyethylene. It was also noted the turnover frequency (TOF) for the Mg₃Al₂O₄ was much greater than the Mg₃Al₂O₃, in agreement with a previous study on supported ethylene polymerisation catalysts which showed this support to be more active.²⁵

However, these supported species displayed much lower activities than the homogeneous catalyst (silica ~70%, Mg₃Al₂O₄ ~8%). The polymethylaluminoxane supported catalyst was by far the most active and selective displaying almost complete selectivity for 1-butene with a TOF almost 7 times that of the homogenous W:MAO = 1:2 (4.0 vs 0.6 mol C₂H₄/mol W⁻¹h⁻¹) after 38 hours at 1 bar ethylene and 100 °C (Fig. 3). This catalyst was also active for selective dimerisation in the gas phase when reacted in an NMR tube with ethylene at 100 °C (Fig. S47).

Supporting of W(NDipp)Me₃Cl on polymethylaluminoxane and subsequent oligomerisation led to complex leaching and an active but inhomogeneous catalyst. This indicates that when W(NDipp)Cl₃(L) is immobilised, the resulting supported species is likely stabilised through a greater number of W-Cl-Al interactions (Scheme 3). Reaction of W(NDipp)Me₃Cl with AlCl₃ results in coordination of the Lewis acid to the halide (Figs S49 & S50) in an analogous fashion to the reactivity reported by Osborn and co-workers for W(NR)(CH₃)₂X (X = F, Cl, Br).³⁸ Such interactions have also been seen in related chromium oligomerisation systems where the chloride ion bound to aluminium can affect the activity of the system.³⁹-⁴¹ Work is currently underway to elucidate the mechanism of the oligomerisation and the role of the polymethylaluminoxane support.

In conclusion, solutions of the complex W(NDipp)Me₃Cl were found to be active for the dimerisation of ethylene in the presence of MAO. Lower cocatalyst ratios result in increased complex stability and hence much higher selectivity for 1-butene. This catalytic activity can be maintained when W(NDipp)Cl₃(THF) is activated on inorganic solids such as a layered double hydroxide, silica or polymethylaluminoxane. The polymethylaluminoxane supported catalyst shows almost complete selectivity for 1-butene with a TOF almost 7 times greater than the homogeneous reaction. This remarkable activity boost may be related to the presence of trimethylaluminium contained within the support, and its apparent porosity.
Experimental

W(NDipp)Cl(L) (L = THF or Et₂O) and polymethylaluminoxane were synthesised according to literature procedures. Layered Double Hydroxides (LDHs) of the type Mg₃AlX where X = CO₃, NO₃ and SO₄ were prepared by the aqueous miscible organic solvent treatment method, and were thermally treated at 150 °C for 6 h under vacuum prior to use.

W(NDipp)MeCl (Dipp = 2,6-Pr₂C₅H₃) W(NDipp)Cl₂ (500 mg, 0.87 mmol) and trimethylaluminium (TMA) (62.7 mg, 0.87 mmol) were dissolved in benzene (~20 mL) and the TMA solution added dropwise to the W(NDipp)Cl₂ solution. The solvent was reduced in vacuo and the resulting brown solid was dissolved in hexane, filtered, reduced to minimum volume and cooled to -30 °C, resulting in orange crystals (300 mg) that were suitable for a single crystal X-ray diffraction study.

Acknowledgements

The authors would like to thank Dr Nicholas H. Rees for solid state NMR spectroscopy, Dr Alexander Kilpatrick for synthesising the polymethylaluminoxane support, Dr Thomas Arnold for help with crystallography and the EPSRC (C.M.R.W.) and SCG Chemicals Co. Ltd, Thailand for funding (Z. R. T and J.-C. B.), and Trinity College, Oxford for a Junior Research Fellowship (Z. R. T).

Notes and references


† Dr P. Dyer and co-workers, Department of Chemistry, University of Durham presented a poster on some aspect of solution phase ethylene dimerisation using W(NDipp)MeCl at UK Catalysis Conference, Loughborough, 2015.