Ethyleneglycol Tungsten Complexes of calix[6 and 8]arenes: Synthesis, Characterization and ROP of ε-caprolactone

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ABSTRACT: By varying the reaction conditions, the reaction of [W(eg)₃] (eg = 1,2-ethanediolato) with p-tert-butylcalix[n]areneH₆ (n = 6 or 8) in refluxing toluene affords, following work-up, a number of products which have been fully characterized. From the reaction of p-tert-butylcalix[6]areneH₆ with one or two equivalents of [W(eg)₃], only the oxo-bridged complex {[W(eg)₂(µ-O)p-tert-butylcalix[6]arene]} (I) could be isolated, whereas the use of four equivalents of [W(eg)₃], in the presence of molecular sieves, afforded {[W(eg)₂]₂p-tert-butylcalix[6]areneH₂} · 2MeCN (2); molecules of 2 pack in bi-layers. Under similar conditions, use of one or two equivalents of [W(eg)₃] and p-tert-butylcalix[8]areneH₈ afforded {[W(eg)₂]₂p-tert-butylcalix[8]arene} · MeCN (3) in which each tungsten centre was bound by four calixarene oxygens. By contrast, the small orange prisms resulting from the use of four equivalents of [W(eg)₃] and p-tert-butylcalix[8]areneH₈ were shown by synchrotron radiation to be a mixture of two isomers (4a/4b · 3.5MeCN). In the major isomer {1,2-[W(eg)₂]₂p-tert-butylcalix[8]areneH₂} (4a), two tungsten centres bind to neighbouring sets of phenolate oxygens, whereas in the minor isomer {1,3-[W(eg)₂]₂p-tert-butylcalix[8]areneH₂} (4b), there is a protonated phenolic group between the two pairs of phenolate oxygens bound to tungsten; the major:minor ratio is about 83:17. Use of p-tert-butyltetrahomodioxacalix[6]areneH₆ with two equivalents of [W(eg)₃] resulted in the isolation of {[WO(eg):p-tert-butyltetrahomodioxacalix[6]areneH₂]} (5 · 0.83toluene:MeCN), in which each dimethyleneoxa bridge is bound to an oxotungsten(VI) centre. Complexes 1 – 5, together with the known complex [W(eg)p-tert-butylcalix[4]arene] (6), have been screened for their ability to ring open polymerize (ROP) ε-caprolactone; for 1, 2 and 6 conversion rates were good (> 88 %) at 110 °C over 12 or 24 h, whereas the calix[8]arene complexes 3 and 4 under the same conditions were inactive.

Introduction

The larger calix[n]arenes (n > 4) continue to be of interest, primarily due to their ability to accommodate multiple metal centres simultaneously.¹ This is desirable since metals in close proximity have the potential to communicate, which in turn may lead to beneficial effects in areas such as magnetism² or catalysis.²b However, the controlled synthesis of specific polymetallic calix[n]arenes can be problematic, often complicated by alkali-metal incorporation or fortuitous hydrolysis and/or oxygenation reactions.³

In the case of tungsten (WCl₅), we have previously shown that by varying the reaction stoichiometry, it is possible to control the degree of metallation.⁴

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The only other tungsten calix[n]arenes (n > 4) that have been reported are the oxo species \([\text{WOCI}_2]_2p\text{-tert-butylcalix}[6\text{areneH}_2]\) and \([\text{WO(NCMe)}_2]_2p\text{-tert-butylcalix}[8\text{arene}]\), the hydrazido complex \([\text{W(NNPh}_2)]_2p\text{-tert-butylcalix}[6\text{arene}]\) plus the \(p\text{-tert-butylcalix}[5\text{arene}]\) complexes prepared by Lattmann et al.\(^7\)

In terms of the ring opening polymerization (ROP) of \(\epsilon\)-caprolactone, few group VI systems appear to have been reported.\(^8\)\(^-\)\(^10\)

Herein, we report a number of new tungstocalix[n]arene systems (see scheme 1), resulting from the reaction of \([\text{W(eg)}_3]\) (eg = 1,2-ethanediolato) and \(p\text{-tert-butylcalix}[6\text{and 8\text{areneH}_6}]\) or \(p\text{-tert-butyltetrahomodiioxacalix}[6\text{areneH}_6].\) The new tungstocalix[6 and 8\text{arene}] have been fully characterized, and their potential as catalysts for the ROP of \(\epsilon\)-caprolactone has been evaluated. We also note that the coordination chemistry of oxacalixarene-type ligands is particularly scant.\(^9\)

Results and Discussion

\(p\text{-tert-butylcalix}[6\text{areneH}_6]\) chemistry

Reaction of \(p\text{-tert-butylcalix}[6\text{areneH}_6]\) with one or two equivalents of \([\text{W(eg)}_3]\) in refluxing toluene afforded, after extraction into hot acetonitrile, the orange/red complex \([\text{[W(eg)]_2(µ-O)}p\text{-tert-butylcalix}[6\text{arene}]}\) (1). The isolated yield of 1, though still only moderate, was greater when two equivalents of \([\text{W(eg)}_3]\) were employed, i.e. ca. 35 % versus ca. 20 % (for one equivalent). In the IR spectrum of 1, the band at 966 cm\(^{-1}\) is tentatively assigned to the bridging oxo group; the latter presumably arises via fortuitous hydrolysis. The \(^1\)H NMR spectrum is consistent with the solid-state structure, vide infra.

Crystals of 1 suitable for an X-ray structure determination were grown from a hot, saturated solution of acetonitrile on slow cooling and prolonged standing at ambient temperature. The compound crystallizes in the non-centric space group \(C2\) with the two halves of the compounds related by a two-fold rotation. Two views of the molecular structure are shown in Figure 1, with selected bond lengths and angles given in the caption. Each tungsten centre adopts a distorted octahedral geometry and these are linked via a near linear oxo bridge \([W(1) – O – W(1)' = 164.7(6)°]\); \(W(1) – O(4) = 1.8953(14) \text{ Å}\), symmetry operator \(i = –x, y, –z\). Each tungsten is further bound by three calix[6\text{arene}] phenoxide groups and a bi-dentate 1,2-ethanediolato group. The conformation of the calixarene ring is best described as an enlarged cup, with the phenolic rings bearing \(O(3)\) and \(O(6)\) forming the base. One of the two molecules of acetonitrile is encapsulated within the cup. Molecules of 2 pack in bi-layers with adjacent \(p\text{-tert-butyl\) groups at one interface and inter-digitated eg groups at the other (see Figure 3).

For each metal, one of the 1,2-ethanediolato ligands is involved in H-bonding to one of the remaining calixarene phenolic groups.

Figure 1 Two views to emphasize the calix[6\text{arene}] conformation in 1. Selected bond lengths (Å) and angles (°): \(W(1) – O(1) 1.873(7), W(1) – O(2) 1.943(4), W(1) – O(3) 1.877(5), W(1) – O(4) 1.8953(14), W(1) – O(5) 1.899(7), W(1) – O(6) 1.855(6); O(1) – W(1) – O(2) 86.9(3), O(4) – W(1) – O(5) 94.3(3), W(1) – O(1) – W(1)’ 164.7(6). (Symmetry operator \(i = –x, y, –z\))

The conformation of the calix[6\text{arene}] ligand is best described as an enlarged cup, with the phenolic rings bearing \(O(3)\) and \(O(6)\) forming the base. One of the two molecules of acetonitrile is encapsulated within the cup. Molecules of 2 pack in bi-layers with adjacent \(p\text{-tert-butyl\) groups at one interface and inter-digitated eg groups at the other (see Figure 3).

Figure 2. Selected bond lengths (Å) and angles (°) for 2:2MeCN:
\[W(1) – O(1) 1.847(6), W(1) – O(2) 1.913(6), W(2) – O(4) 1.855(5), W(2) – O(5) 1.922(5); O(1) – W(1) – O(2) 86.0(3), O(7) – W(1) – O(10) 163.5(3), O(4) – W(2) – O(5) 86.5(2), O(11) –

\[W(1) – O(1) 1.873(7), W(1) – O(2) 1.943(4), W(1) – O(3) 1.877(5), W(1) – O(4) 1.8953(14), W(1) – O(5) 1.899(7), W(1) – O(6) 1.855(6); O(1) – W(1) – O(2) 86.9(3), O(4) – W(1) – O(5) 94.3(3), W(1) –

\[W(1) – O(1) – W(1)’ 164.7(6). (Symmetry operator \(i = –x, y, –z\))

The conformation of the calix[6\text{arene}] ligand is best described as an enlarged cup, with the phenolic rings bearing \(O(3)\) and \(O(6)\) forming the base. One of the two molecules of acetonitrile is encapsulated within the cup. Molecules of 2 pack in bi-layers with adjacent \(p\text{-tert-butyl\) groups at one interface and inter-digitated eg groups at the other (see Figure 3).
W(2) – O(14) 164.3(2).

Figure 3. Bi-layer packing of 2·2MeCN.

p-tert-butylcalix[8]areneH₈ chemistry

On extending the synthetic methodology to p-tert-butylcalix[8]areneH₈, use of one or two equivalents of [W(eg)₃] led to the isolation of the red complex [W(eg)]₂p-tert-butylcalix[8]arene (3) in good isolated yield (ca. 60 % for two equivalents and ca. 40 % for one equivalent).

Large crystals (see ESI, Figure S1) of 3·MeCN suitable for an X-ray structure determination were grown from a hot, saturated solution of acetonitrile on slow cooling and prolonged standing at ambient temperature. The asymmetric unit comprises only one molecule of 3. The molecular structure is shown in Figure 4, with selected bond lengths and angles given in the caption.

Figure 4. Molecular structure of compound 3·MeCN. Selected bond lengths (Å) and angles (°): W(1) – O(1) 1.908(3), W(1) – O(2) 1.864(3), W(1) – O(3) 1.954(3), W(1) – O(4) 1.861(3), W(2) – O(5) 1.883(2), W(2) – O(6) 1.883(3); W(1) – O(1) – C(1) 1.34.5(2), O(1) – W(1) – O(3) 171.89(11), O(2) – W(1) – O(4) 101.06(12).

The p-tert-butylcalix[8]arene is fully deprotonated and adopts a twisted conformation to accommodate, on opposite sides of the molecule, the binding of the two pseudo-octahedral tungsten centres. Each tungsten centre is an average 0.955(7) Å out of the plane of the calix[8]arene ring. Similar conformations were reported for the complexes [{WO(NCMe)}₂p-tert-butylcalix[8]arene] and [{MCl₂}₂p-tert-butylcalix[8]arene} (M = Nb or W).

Increasing the amount of [W(eg)] (to four equivalents) resulted in the formation of an orange crystalline solid, however the ¹H NMR spectrum is somewhat different (more complicated) to that of 3. Interestingly, a crystal structure determination, requiring the use of synchrotron radiation, revealed that the product was a mixture of isomers. In the major isomer (Figure 5, top), the two tungsten centres bind to neighbouring pairs of phenolate oxygens. For this isomer, all phenolic protons (at O(5), O(6), O(7) and O(8)) were located, but not for O(2) in the minor component (Figure 5, bottom).

Figure 5. Top, molecular structure of the major isomer 4a; bottom, molecular structure of minor isomer 4b. Selected bond lengths (Å) and angles (°): Major: W(1) – O(1) 1.954(4), W(1) – O(2) 1.904(4), W(2) – O(3) 1.912(4), W(2) – O(4) 1.761(6), W(1) – O(9) 1.896(5), W(1) – O(10) 1.942(5); O(1) – W(1) – O(2) 89.58(19), O(1) – W(1) – O(9) 159.5(2), O(3) – W(2) – O(4) 89.3(2). Minor: W(1X) – O(1) 1.855(4), W(2X) – O(3) 1.861(4), W(2X) – O(4) 2.242(7), W(1X) – O(8) 2.150(5), W(1X) – O(9X) 1.88(2), W(1X) – O(10X) 1.88(2); O(1) – W(1X) – O(9X) 104.9(8), O(1) – W(1X) – O(9X) 78.8(6), O(3) – W(2X) – O(4) 77.4(2).

In the minor isomer, this phenolic group O(2) resides between
pairs of bonding phenolate oxygens. The tungsten centres and the eg ligands are disordered such that W(1):W(1X) = 83.3:16.6(8)% and W(2):W(2X) = 62.6:37.4(5)% (same occupancy for the attached eg ligands). The W(1):W(1X) gives the isomer ratio, whereas W(2):W(2X) is more of a conformational issue, probably partially depending on the location of W(1) or W(1X).

The acetonitrile containing N(1) lies in the cleft between phenols containing O(5) and O(6). That containing N(3) lies between O(1) and O(2), whilst those containing N(2) and N(4) lie exo to the calixarene, between complexes.

p-tert-butyltetraphomiodoxacalix[6]areneHs chemistry

Reaction of p-tert-butyltetraphomiodoxacalix[6]areneHs with two equivalents of [W(eg)p-

acetonitrile containing N(1) lies in the cleft between phenols containing O(5) and O(6). That containing N(3) lies between O(1) and O(2), whilst those containing N(2) and N(4) lie exo to the calixarene, between complexes.

Figure 6. View of the molecular structure of 5. Selected bond lengths (Å) and angles (°): W(1) – O(1) 1.906(4), W(1) – O(2) 1.891(4), W(1) – O(7) 1.908(4), W(1) – O(11) 2.394(4), W(2) – O(4) 1.941(4), W(2) – O(5) 1.879(4), W(2) – O(8) 1.890(4), W(2) – O(45) 2.403(4); O(7) – W(1) – O(11) 172.46(19), O(8) – W(2) – O(45) 174.21(19).

Ring opening polymerization screening

The complexes 1 – 5 have been screened for their ability to ring open polymerize ε-caprolactone in the presence of benzyl alcohol (BnOH). For comparison, the known complex [W(eg)p-


It was necessary to activate 2 for the ROP of ε-caprolactone using BnOH. Good conversions were only achieved at temperatures over 100 °C, over prolonged periods (> 12 h), e.g. see runs 1 – 3 versus 4 - 6 (Table 1). Based on entries 1 to 5, there is a near linear relationship between monomer conversion and number average molecular weight (Mn), with narrow molecular weight distributions (≤1.34). Moreover, a linear relationship between [CL]/[W] and Mn demonstrates the classical feature of a living process (see Fig. 7). On increasing the temperature beyond 110 °C, the conversion rates and average molecular weight values (Mn) decreased, suggesting that 110 °C is more favourable for catalyst stability. Increasing the molar ratio of ε-CL:Al from 100:1 to 1000:1 (runs 13 - 16, Table 1) led to an increase in the observed molecular weights 18.82 × 10^3 to 52.83 × 10^3, which is a desirable scenario given higher molecular weight polyesters possess better mechanical properties. However, although there was little change in the molecular weight distribution (1.36 – 1.83), the conversion rate dropped considerably (from 99% to 56%).

Using the optimized conditions for 2, we also screened the complexes prepared herein as well as the known [W(eg)p-tert-butyldcalix[4]arene] (6). Results in Table 2 revealed good conversions for those systems derived from calix[6]arene ligands (1 and 2), although unlike for ethylene polymerization, we did not observe any beneficial effects (versus methylene, –CH2–) herein when using the dioxaamethylene (–CH2OCH2–) bridged complex 5. Surprisingly, the systems 3 and 4 derived from calix[8]arene ligands were inactive under similar conditions. The reason for this is not clear, but may be related to the increased conformational flexibility of the calix[8]arene ligand which somehow hinders the ring opening polymerization process. Furthermore, the known complex [W(eg)p-tert-butyldcalix[4]arene] (6) was found to be active, affording a conversion rate of 74%, somewhat lower that observed for the calix[6]arene-based systems.

In terms of polymer characterization (molecular weight determination and end group analysis), 1H and 13C NMR spectra of selected polymers (entries 1, 2 and 5, Table 2) were recorded.
(see ESI, Figures S3 – S10). Signals at around δ 7.37 and 5.15 ppm (C6H1:CH2-), and 3.67 ppm (CH2:CH2:OH), with an intergral ratio of 5:2:2, indicated that the polymer chains are capped by a benzyl ester group and a hydroxy end group. This suggests that the polymerization occurs through insertion of a benzyl alkoxy group into CL. The MALDI-TOF spectra (ESI, Figures S11 - S13) revealed the presence of the benzyloxy initiating group and a series of peaks separated by 114.14 mass units.

In summary, a number of new tungstocalix[6 and 8]arenes have been prepared using the metal precursor [W(eg):3] (eg = 1,2-ethanediol) and the calix[n]arenes p-tert-butylicalix[6 and 8]areneH1.8 or p-tert-butyltetrahydrodioxacalix[6]areneHs. Crystal structure determinations reveal a preference for binding two metal centres, though in the solid-state, the relative positions of the metal centres at the lower rim of the calixarene can vary, for example 1,2-[W(eg):2] versus 1,3-[W(eg):2]:p-tert-butylicalix[8]areneHs. The syntheses are sensitive to the presence of air and/or moisture, and the work herein includes the characterization of two oxotungsten species. In terms of the ROP of ε-caprolactone, only negligible polymer was isolated at prolonged standing (2 - 3 days) at ambient temperature afforded red prisms. Yield 0.75 g, 35.4 %. Elemental analysis calculated for 1, C70H100.5N1.5O14W2, C62H113.5N0.5O12W2 C 62.1, H 6.4 , N 0.4 % ; Found C 61.3, H 5.6 %; Found C 57.2, H 6.5, N 1.1 %. IR (nujol mull, KBr): 1596w, 1297m, 1260s, 1208s, 1118s, 1106s, 1071s, 1051s, 795w, 771m, 723s, 678w, 640s, 616s, 585m, 567m, 494w. Mass Spec (EI): 1207 (M + – W – 3egH2). 1H NMR (CD3CN, 600 MHz) δ : 7.46 (s, 8H, arylH), 7.14 (s, 4H, arylH), 5.90 (s, 8H, OCH2), 5.05 (d, 3JHH = 15.2 Hz, 2H, endo-CH2), 4.81 (d, 3JHH = 13.2 Hz, 4H, endo-CH2-), 3.51 (d, 3JHH = 13.2 Hz, 4H, exo-CH2), 3.39 (d, 3JHH = 15.2 Hz, 2H, exo-CH2), 2.17 (s, 12H, 4MeCN), 1.35 (s, 36H, C(CH3)3)), 1.18 (s, 18H, C(CH3)3)).

**Experimental**

**General:** All manipulations were carried out under an atmosphere of dry nitrogen using conventional Schlenk and cannula techniques or in a conventional nitrogen-filled glove box. Diethyl ether and tetrahydrofuran were refluxed over sodium and benzophenone. Toluene was refluxed over sodium. Dichloromethane and acetonitrile were refluxed over calcium hydride. All solvents were distilled and degassed prior to use. IR spectra (nujol mulls, KBr windows) were recorded on a Nicolet Avatar 360 FT IR spectrometer. 1H NMR spectra were recorded at room temperature on a Varian VX4 400 S spectrometer at 400 MHz or a Gemini 300 NMR spectrometer or a Bruker Advance DPX-300 spectrometer at 300 MHz. The 1H NMR spectra were calibrated against the residual protio impurity of the deuterated solvent. Elemental analyses were performed by the elemental analysis service at the London Metropolitan University and at the University of Hull. Matrix Assisted Laser Desorption/Ionization Time of Flight (MALDI-TOF) mass spectrometry was performed in a Bruker autoflex III smart beam in linear mode, and the spectra were acquired by averaging at least 100 laser shots. 2,5-Dihydroxybenzoic acid was used as the matrix and THF as solvent. Sodium chloride was dissolved in methanol and used as the ionizing agent. Samples were prepared by mixing 20 μl of matrix solution (2 mg/ml) with 20 μl of matrix solution (10 mg/ml) and 1 μl of a solution of ionizing agent (1 mg/ml). Then 1 ml of these mixtures was deposited on a target plate and allowed to dry in air at ambient temperature. The ligands p-tert-butylicalix[6]areneH6, p-tert-butylicalix[8]areneH8 and p-tert-butyltetrahydrodioxacalix[6]areneHs were prepared as described in the literature. The precursor [W(eg):1] and complex 6 were prepared via the methods of Lehtonen and Sillanpää.12,14


Toluene (30 ml was added to a Schlenk containing p-tert-butylicalix[6]areneH6 (1.40 g, 1.44 mmol) and [W(eg):1] (1.40 g, 2.91 mmol). After refluxing for 12 h, the volatiles were removed in-vacuo, and the residue was extracted into hot (brought to reflux with a heat gun) acetonitrile (30 ml). Prolonged standing (2 - 3 days) at ambient temperature afforded red prisms. Yield 0.75 g, 35.4 %. Elemental analysis calculated for 1, C70H100.5N1.5O14W2 C 62.1, H 6.4 , N 0.4 %; Found C 61.3, H 5.6 %; Found C 57.2, H 6.5, N 1.1 %. IR (njoulull, KBr): 1596w, 1297m, 1260s, 1208s, 1118s, 1106s, 1071s, 1051s, 795w, 771m, 723s, 678w, 640s, 616s, 585m, 567m, 494w. Mass Spec (EI): 1207 (M + – W – 3egH2). 1H NMR (CD3CN, 400 MHz) δ : 7.46 (s, 8H, arylH), 7.14 (s, 4H, arylH), 5.90 (s, 8H, OCH2), 5.05 (d, 3JHH = 15.2 Hz, 2H, endo-CH2), 4.81 (d, 3JHH = 13.2 Hz, 4H, endo-CH2-), 3.51 (d, 3JHH = 13.2 Hz, 4H, exo-CH2), 3.39 (d, 3JHH = 15.2 Hz, 2H, exo-CH2), 2.17 (s, 12H, 4MeCN), 1.35 (s, 36H, C(CH3)3)), 1.18 (s, 18H, C(CH3)3)).


As for 1, but using p-tert-butylicalix[6]areneH6 (1.40 g, 1.44 mmol) and [W(eg):1] (1.40 g, 2.91 mmol) and activated molecular sieves. Yield 1.06 g, 44.4 %. Elemental analysis calculated for 2·MeCN (sample dried in vacuo – MeCN), C77H106.5N1.5O14W2 C 56.4, H 6.2, N 0.9 %; Found C 56.7, H 6.5, N 1.1 %. IR (njuolull, KBr): 1639 w, 1303 s, 1261 s, 1201 m, 1155 m, 1076 m, 1050 m, 966 m, 936 m, 914 m, 888 m, 857 w, 795 w, 771 m, 723 s, 678 m, 648 w. Mass Spec (EI): 1468 (MH+). 1H NMR (CD3CN, 400 MHz) δ : 7.46 (s, 8H, arylH), 7.14 (s, 4H, arylH), 5.90 (s, 8H, OCH2), 5.05 (d, 3JHH = 15.2 Hz, 2H, endo-CH2), 4.81 (d, 3JHH = 13.2 Hz, 4H, endo-CH2-), 3.51 (d, 3JHH = 13.2 Hz, 4H, exo-CH2), 3.39 (d, 3JHH = 15.2 Hz, 2H, exo-CH2), 2.17 (s, 12H, 4MeCN), 1.35 (s, 36H, C(CH3)3)), 1.18 (s, 18H, C(CH3)3)).
Table 1. Ring Opening Polymerization of ε-CL by pre-catalyst 2.

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*Conditions: 20 μmol of cat.; 1.0 M ε-CL toluene solution. ^X=W complex *GPC data in THF vs polystyrene standards.

Preparation of (1,2-[W(eg)3]2-p-tert-butyltetrahydroxaloxalix[6]areneH6) (5)

4.91 (m, 2H, endo-CH2), 4.47 (overlapping d, 4H, endo-CH2),
4.10 (m, 2H, endo-CH2), 3.46 (overlapping d, 4H, exo-CH2),
3.29 (m, 2H, exo-CH2), 2.91 (m, 2H, exo-CH2), 2.17 (s, 10.5H.
MeCN), 1.32 (s, 36H, C(CH3)3), 1.31 (s, 18H, C(CH3)3), 1.22 (s,
18H, C(CH3)3).

As for 1, but using p-tert-butyltetrahydroxaloxalix[6]areneH6
(0.50 g, 0.49 mmol) and [W(eg)3] (0.36 g, 1.01 mmol). Yield
0.56 g, 69.1 %. Elemental analysis calculated for 5,
C72H80O4W2 (sample dried in vacuo for 12 h): C 55.9, H, 6.0 %;
Found C 55.6, H 6.1 %. IR (nujol mull, KBr): 3170 bw, 1635 w,
1597 w, 1304 s, 1155 m, 1077 m, 965 m, 892 m, 846 w, 770 m, 723 s.

Preparation of [1,2-[W(eg)2]2-p-tert-butylcalix[8]areneH4]/(1,2-
[W(eg)2]2-p-tert-butylcalix[8]areneH4) (4a)/(4b) 5.35 MeCN
As for 2, but using p-tert-butylcalix[8]areneH4 (1.00 g, 0.78 mmol)
and [W(eg)2] (1.11 g, 3.10 mmol). Yield 1.27 g, 79.4 %.

 Element analysis calculated for 4a/b 5.35 MeCN,
C105H134N3.5O16W2 C 60.5; H 6.6; N 2.4 %; Found C 60.4, H
6.7, N 2.4 %. IR (nujol mull, KBr): 3183, 1597 w, 1297 m,
1254 s, 1208 s, 1198 s, 1107 m, 1070 m, 1051 m, 945 m,
915 m, 865 m, 830 m, 803 m, 796 m, 760 m, 723 s, 678 w,
641 s, 616 w. Mass Spec (EI): 1922 (MH+ – 3MeCN),
1777 (MH+ – 3.5MeCN – 2eg), 1715 (MH+ – 3.5MeCN – 3eg),
1657 (MH+ – 0.5MeCN – W(eg)3). 1H NMR (C6D6, 500 MHz)
δ: 7.52 (m, 2H, arylH), 7.16 (m, 2H, arylH), 7.00 (m, 2H, arylH),
5.63 (m, 4H, OCH2), 5.29 (m, 4H, OCH2), 5.14 (m, 4H, OCH2),
5.43 (m, 4H, OCH2), 5.29 (m, 4H, OCH2), 5.14 (m, 4H, OCH2),
5.43 (m, 4H, OCH2), 5.29 (m, 4H, OCH2), 5.14 (m, 4H, OCH2),
5.43 (m, 4H, OCH2), 5.29 (m, 4H, OCH2), 5.14 (m, 4H, OCH2).
Crystallography

For 1, 3, and 5: Data were collected using an Agilent Xcalibur diffractometer with an Eos CCD detector operating with Mo Kα radiation using a series of ω-scans. Data were treated using an empirical absorption correction. For 1 and 5 data collection was conducted at room temperature (293 K). For 3 data were collected with the crystal cooled to 135 K. Structure solution and refinement was conducted using the SHELX suite of programs.16

Structure of 1: Two of the three symmetry independent tert-butyl groups (C9-C11 and C20-C22) were modelled as disordered over two sets of positions. There was no evidence for inclusion of solvent within the structure. Structure of 3: No disorder was modelled for the tert-butyl groups although these exhibit relatively large displacement parameters, but the core of the calix[8]arene is well determined. Small portions of the structure contained disordered solvent molecules which could not be resolved as point atoms. The Platon SQUEEZE routine17 was employed to deal with scattering from these regions. For 1 and 3 the unit cell contained four disordered MeCN molecules. Structure of 5: The central core of the oxacalixarene is well determined but the tert-butyl groups show rather larger displacement parameters. The structure contains a single molecule of MeCN within each oxacalixarene bowl and a molecule of toluene (~83 % occupied) that is located on the bottom edge of the oxacalixarene and may be localised by C–H...π interactions.

For 2: Data collected on a Bruker APEX 2 CCD diffractometer at Daresbury SRS station 9.8.18 The tert-butyl groups at C(18) and C(40) were modelled as disordered over two sets of positions. The MeCN in the calixarene cavity was well defined and was modelled as point atoms. Platon SQUEEZE17 was used to model eight other MeCN molecules per unit cell (one per complex molecule) as diffuse electron density. The structure could also be modelled in the triclinic crystal system, but this led to two very large residual electron density peaks of > 8 eÅ³, so the simpler C2/c solution was deemed more satisfactory. For 4: Data collected on a Bruker APEX 2 CCD diffractometer at Daresbury SRS station 9.8.19 Both W(1) and W(2) atoms were modelled as two-fold disordered along with the attached eg groups, with occupancy factors of 83.39:16.61(8)% for the former, and 62.6:37.4(5)% for the latter. The tert-butyl group at C(18) was modelled with all atoms disordered over two sets of positions, while those at C(40) and C(51) were modelled with just the methyl groups split over two sets of positions. One MeCN was 50/50 disordered over a centre of symmetry.

Crystal structure data have been deposited with the CCDC (numbers 989342-989344, 989597, and 1010285). These data can be obtained from of charge from www.ccdc.cam.ac.uk.

Acknowledgments

We thank Sichuan Normal University for financial support. The CCLRC is thanked for the award of beamtime at SRS Daresbury Laboratory (Station 9.8), and the EPSRC Mass Spectrometry Service at Swansea is thanked for data.

References


Table 2. Ring Opening Polymerization of ε-CL by pre-catalysts 1 - 6.

<table>
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<th>Entry</th>
<th>Cat.</th>
<th>CL:X_b:BuOH</th>
<th>T/°C</th>
<th>r/h</th>
<th>m/g</th>
<th>Yield (%)</th>
<th>Mw*10^3 c</th>
<th>PDI</th>
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*Conditions: 20 μmol of cat.; 1.0 M ε-CL toluene solution. 

CCL toluene solution.

*X=W complex ‘GPC data in THF vs polystyrene standards.
Table 3. Crystallographic data for structures 1 - 5

<table>
<thead>
<tr>
<th>Compound</th>
<th>C₇₀H₈₂O₁₁W₂</th>
<th>C₇₄H₉₆O₁₄W₂·2MeCN</th>
<th>C₇₂H₁₁₂O₁₂W₂·3.5 MeCN</th>
<th>C₹₂H₁₂₄O₁₆W₂·0.83PhMe MeCN</th>
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<tr>
<td>Formula</td>
<td>C₇₀H₈₂O₁₁W₂</td>
<td>C₇₄H₉₆O₁₄W₂·2MeCN</td>
<td>C₇₂H₁₁₂O₁₂W₂·3.5 MeCN</td>
<td>C₹₂H₁₂₄O₁₆W₂·0.83PhMe MeCN</td>
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<td>Formula weight (g mol⁻¹)</td>
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Transmission factors (max., min) | 0.610, 0.503 | 0.924, 0.617 | 0.772, 0.611 | 0.933, 0.679 |
Crystal size (mm³) | 0.29 × 0.24 × 0.22 | 0.20 × 0.09 × 0.03 | 0.20 × 0.15 × 0.10 | 0.18 × 0.08 × 0.03 |
Reflections measured | 6695 | 95194 | 17552 | 60220 |
Unique reflections, Rint | 6695 | 27094 | 17552 | 31223 |
R ws | 0.0258 | 0.1768 | 0.0536 | 0.0443 |
Reflections with F² > 2σ(F²) | 5824 | 22156 | 14374 | 20519 |
Number of parameters | 362 | 899 | 979 | 1435 |
R₁, wR² [F² > 2σ(F²)] | 0.0364, 0.0841 | 0.0889, 0.1962 | 0.0387, 0.0738 | 0.0763, 0.1588 |
GOOF | 1.02 | 1.22 | 1.07 | 1.11 |
Largest difference peak and hole (e Å⁻³) | 1.296 and -0.645 | 2.734 and -2.488 | 1.329 and -0.872 | 1.318 and -2.295 |

GOOF: Goodness of fit; R₁: R₁ = ∑|Fobs - Fcalc|/∑|Fobs|; wR²: wR² = ∑[(wF²o-F²c)²]/∑[(wF²o)²]; R₁ and wR² are calculated for reflections with F² > 2σ(F²).