Grafted Metallocalixarenes as Single-Site Surface Organometallic Catalysts

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Abstract: Metallocalixarenes were grafted onto silica using a surface organometallic approach and shown to be active and selective catalysts for epoxidation of alkenes using organic hydroperoxides. Calixarene--TiV precursors were anchored at surface densities from 0.1 to near-monolayer coverages (0.025–0.25 calixarene nm−2). Several spectroscopic methods independently detected calixarene--TiV connectivity before and after epoxidation catalysis. Kinetic analyses of cyclohexene epoxidation confirmed that the active sites were anchored on the silica surface and were significantly more active than their homogeneous analogues. The steric bulk and multidentate binding of the calixarenes led to structural stability and to single-site behavior during epoxidation catalysis. Rate constants were independent of surface density for cyclohexene epoxidation with tert-butyl hydroperoxide (11.1 ± 0.3 M−1 s−1) or cumene hydroperoxide (25 ± 2 M−1 s−1). The materials and methods reported here allow the assembly of robust surface organometallic structures in which the active sites behave as isolated species, even near saturation monolayer coverages. In turn, this makes possible the rational design and synthesis of a class of heterogeneous oxide catalysts with atomic-scale precision at the active site.

Introduction

Many heterogeneous catalysts currently in practice are chemically complex and consist of oligomeric active site structures that are difficult to optimize via atomic-scale control at the active site. A promising new method for the design and synthesis of heterogeneous catalysts with control of active site structure involves the use of emerging concepts in surface organometallic chemistry. Surface organometallic catalysts consist of a transition-metal-containing active site that is grafted onto a solid support, which serves as a rigid ligand for the metal center.1–5 These catalysts can exhibit interesting support effects, including large rate enhancements, when active sites are anchored onto supports instead of acting as homogeneous entities in solution.5,6 These rate enhancements resemble those reported for systems consisting of hybrid organic–inorganic materials,6 and reflect the synergy between the catalytically active site and a support surface that acts as a ligand.

We report here a new strategy to control the interplay between metal centers and organic ligands or surfaces in grafted organometallic catalysts. Our approach involves confining an isolated metal atom within a pseudotetrahedral oxo coordination environment, defined from the top by a calixarene molecule and from the bottom by a silica surface. The rigidity and steric bulk of both ligands—calixarene and silica—ensure metal atom site isolation, even during kinetic processes that require ligand exchange; such stability is essential for durable and robust heterogeneous catalysts.

Our synthesis of grafted metallocalixarene catalysts is based on anchored calixarene structures 1 and 2. These have been synthesized according to previously reported procedures by treating a calixarenetriol or -tetrol compound with an activated silica containing chloride ligands.7 We previously showed that this process leads to the grafting of a calixarene monolayer and to a 10 Å decrease in the average pore radius of the silica support, consistent with the expected thickness of a single calixarene layer. Each calixarene in 1 was shown to act as a stoichiometric adsorbent for one toluene molecule, a result that is consistent with an open cone conformer for each anchored calixarene.7

Koiland molecules8 originally inspired the synthesis of grafted calixarenes 1 and 2. Reports of isostructural Ti dimers of calixarenes,9 as well as structures in which calixarenes serve as well-defined oxo surface for several other metals,10 suggested to us that the Si atom in 2 could be successfully replaced by transition metals of catalytic interest. We report

Our objective in substituting Ti within the grafted metallocalixarene architecture is to incorporate epoxidation catalytic activity in the same manner as for titanium silicalite (TS-1), in which isolated Ti atoms replace Si atoms within the zeolite framework.12–14 We demonstrate the utility of 3 as a highly active, stable, and selective catalyst for alkene epoxidation. Grafted titanium complexes catalyze epoxidation reactions but lose activity and selectivity during catalysis, especially at high temperature.2,4,18

### Experimental Section

**Precursor Synthesis.** Calixarene–Ti⁴⁺Cl⁴⁻ 4b was prepared by refluxing 1.0 equiv of dimethoxyxcalixarene 5a (Acros, 99%) with TiCl₄ in toluene (Aldrich, 1.0 M) for 48 h under reflux by recrystallization from hexane according to previously published procedures.11 Elemental analysis (Anal. Calcd for 4b(CH₃)₃O, C: 73.79; H: 7.55. Found: C: 73.97; H: 7.82) and ¹H NMR (400 MHz, CDCl₃, 298 K) (δ 1.21 (m, 36H, Bu'), 3.34 (dd, 4H, J = 12.4 Hz, exo-CH₂O), 4.23 (s, 3H, OCH₃), 4.35 (d, 2H, J = 12.4 Hz, endo-CH₂O), 4.78 (d, 2H, J = 12.4 Hz, endo-CH₂), 7.05 (m, 8H, ArH)) agreed with published values. The four doublets in the methylene region are characteristic of the lower symmetry of the mono-demethylated product 4b. Calixarene 5a has only two methylene doublets. Species 4b could be stored for several weeks as a dark red toluene solution under Schlenk conditions without detectable degradation.

**Immobilization of 4b To Yield 3-75.** In the naming scheme used here, 3-75 indicates a material with 75 μmol of calixarene–Ti⁴⁺ per g of catalyst. Chromatography silica gel (300 mg, 60 Å pores, 250–500 μm, Selecto) was partially dehydroxylated under dynamic vacuum at 500 °C for 24 h. The silica was transferred to a flask with 50 mL of anhydrous toluene. A 25 μmol sample of 4b (as a toluene solution) was added and the suspension refluxed under N₂. The red color of 4b was gradually transferred from the solution to the solid. After 0.5 h, 70 μL of 2,6-di-tert-butylpyridine (0.25 mmol) was added, and the suspension was returned to reflux. After 24 h, the solid was filtered, washed with ~300 mL of hot anhydrous toluene, and dried under dynamic vacuum at 25 °C for at least 4 h and at 250 °C for 1 h. Materials with identical catalytic behavior were made with and without recrystallization of 4b, thus enabling the entire catalyst synthesis to occur in a single pot using commercially available reagents.

**Immobilization of 5b To Yield 2-116.** Using our previously published procedures,7 a suspension of 1.00 g of chlorinated silica gel was refluxed in 50 mL of toluene with 160 mg of 5b⁵⁻ (0.24 mmol) and 1.0 mL of N₂ (7.2 mmol, 2 equiv relative to surface silanols) to yield the silicon-containing analogue of 3.

**Catalysis.** A 50 mg sample of catalyst (7 μmol of Ti for 3-138) and ~300 mg of 4A or 3A molecular sieves were added to a 50 mL round-bottom flask with magnetic stirring. The reactor was sealed and degassed under dynamic vacuum at 25 °C for 1 h and at 70 °C for 1 h. The reactor was cooled under Ar backflow, charged with 40 mL of octane (solvent) and 6.5 mmol of substrate, sealed with a PTFE-coated rubber septum, and transferred to a 60 °C oil bath. After thermal equilibration, 1.5 mmol of tert-butyl hydroperoxide (TBHP; Fluka, ~5.5 M solution in nonane with 4A molecular sieves) or cumyl hydroperoxide (CHP; Fluka, ~80% solution in cumene, 4A molecular sieves added) was injected to start the reaction. Aliquots (100 μL) were removed periodically using a syringe, filtered to remove the catalyst, and analyzed using an Agilent 6890 GC system equipped with a flame ionization detector using an HP-1 methylsilicone capillary column. Cumene or nonane present in the hydroperoxide solution was used as an internal standard. Details of reagent purification and of product identification are given in the Supporting Information.

**Analytical Characterization.** Thermogravimetric analysis (TGA) was performed on a TA Instruments TGA 2950 system using a flow of dry synthetic air at a ramp rate of 5 °C/min. Carbon content was measured by the Berkeley Microanalytical Laboratory using a Perkin-Elmer 2400 Series II combustion analyzer. Titanium content was measured by Quantitative Technologies, Inc. using inductively coupled plasma mass spectrometry (ICP-MS).

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Spectroscopy. UV–vis spectroscopy was performed on a Varian Cary 400 Bio UV–vis spectrophotometer equipped with a Harrick Praying Mantis accessory for diffuse reflectance measurements on solids at room temperature. Compressed PTFE powders were used as a reference. Fluorescence spectra were measured using a Hitachi F-4500 fluorescence spectrophotometer with a solids reflectance accessory. Solid-state 13C CP/MAS NMR was performed at the Caltech solid-state NMR facility using a Bruker DSX500 spectrometer operating at 500 MHz.

Results and Discussion

Materials Synthesis and General Characterization. The synthetic scheme for grafting metallocalixarenes is depicted in Scheme 1 and described in the Experimental Section. The catalyst synthesis can be generalized to the large library of upper- and lower-rim substituted calixarenes,23 but care must be exercised to prevent the formation of multinuclear species.9,24,25 Precursor 4b was immobilized with quantitative yield up to a surface density of 0.18 calixarene nm$^{-2}$ (0.15 mmol g$^{-1}$). The surface density was increased to a maximum of 0.25 calixarene nm$^{-2}$ (0.20 mmol g$^{-1}$) when an excess of 4b was refluxed in the presence of silica for 280 h. After this point, additional 4b had no effect on the ultimate surface coverage of 3 determined by TGA and elemental analysis. We note that the silanol density remains much higher than the calixarene active site density for all materials synthesized and is not appreciably changed upon calixarene grafting for silica pretreatment temperatures below 1000 °C.26

A Ti atom:calixarene molecule ratio of 1.0 ± 0.1 was independently measured in 3 at all surface coverages (0.03–0.25 nm$^{-2}$) by combustion analysis and ICP-MS. These equimolar ratios strongly indicate that the molecular precursor 4b was grafted intact. Bulk calixarene contents were estimated by total weight loss in TGA above 250 °C (Supporting Information Figure S1). The onset of weight loss upon heating 3 in air is quite dramatic, with a sharp maximum in the derivative plot centered between 360 and 380 °C. Calixarene


contents were measured from combustion analysis and TGA, and both methods agreed to within 10%. Nitrogen physisorption data and BJH pore size distributions are similar to those reported previously for 1 and show decreasing total pore volume and average pore radius as calixarene loading is increased (Supporting Information Figure S2). The decrease in pore volume corresponds to an average calixarene–TiIV molecular volume of 2.3 nm³, in reasonable agreement with the molecular dimensions predicted from molecular models (1.4 nm x 1.4 nm x 1.0 nm = 2.0 nm³). Additional materials characterization data are provided in Table 1.

Materials Spectroscopic Characterization. NMR spectroscopy provides additional compelling evidence that the atomic connectivity between Ti and calixarene in 3 is retained during grafting. Close comparison of 13C NMR spectra for materials 2 and 3 with the solid state and for molecular precursors 4a28 and 4b27 in deuterated chloroform shows that all four spectra are similar, with the exception of two resonances, 4 and 7. In the spectra of the precursors, resonances 4 and 7 are shifted downfield 8 and 13 ppm, respectively, when Ti-containing 4a is compared to unmetalated 5a. Likewise, in the solid-state spectrum of 3, resonances 4 and 7 are shifted by 2 and 11 ppm, respectively, relative to the resonances in 2 (Figure 1). These shifts are consistent with expected changes in electron density upon phenoxide (resonance 7) and methyl ether (resonance 4, inset) coordination to Ti. The latter coordination represents a weak dative bond inferred from interatomic distances between Ti and methyl ether oxygen in structures derived from single-crystal X-ray diffraction of 4a,27 4b,31 and related titanocalixarenes.9 The resonance assignment for carbon 4 was verified by isotopically enriching the methoxy position in grafted calixarene materials.

UV–vis and fluorescence spectroscopies were used to probe the surface structures formed by grafting of 4b onto silica. The absorption spectra of precursor 4b and of immobilized 3 and 2 are shown in Figure 2A. All materials exhibit an absorption maximum at 270–290 nm arising from π→π* transitions, but only calixarene–TiIV complexes 3 and 4b show an additional broad absorption feature at 290–500 nm, which we assign to calixarene–TiIV charge transfer (LMCT) bands. Material 6 was produced by treatment of 3 in dry air at 50 °C. This process removed all absorption features in the 290–500 nm range, confirming that these electronic transitions arise from the calixarene ligands and also ruling out the presence of extended Ti–O–Ti connectivity even after calcination. Fluorescence emission spectra (λex = 270 nm; Supporting Information Figure S3) of 3 show strong quenching relative to those of both the unmetalated calixarene 2 and the calcined 6. This quenching is a common feature of d⁰ metal–dye complexes,30 and is consistent with Ti attachment to the calixarene in 3.

We use the position of the calixarene–TiIV LMCT edge as a probe of catalyst structure, in much the same way as edge energies are used to characterize dispersed metal oxides.31,32 As seen in Figure 2B, extrapolation of the linear region to the intercept gives the same edge energy of 2.18 ± 0.02 eV (or 585 nm) for all grafted materials, irrespective of titanium loading. This strongly suggests that the structure of the predominant grafted species is independent of calixarene–TiIV surface density. In dispersed metal oxides, the edge energy is unchanged at low surface density, because of the predominant presence of isolated o xo species, and then decreases as metal centers become linked by bridging oxygens with increasing surface coverage.31,32 We attribute the constant edge energy in 3 up to the maximum surface loading possible to structural isolation enforced by the steric bulk of the calixarene and silica surface as ligands. The edge energy of the precursor molecule 4b lies at 1.90 eV, 0.28 eV lower than that of the grafted 3, reflecting the change of ligand from R₃SiO⁻ in 3 to Cl⁻ in 4b and confirming that the Ti precursor has been covalently grafted, instead of merely physisorbed, on the surface.

![Figure 1. 13C CP/MAS NMR spectra of immobilized materials 2 (M = Si) and 3 (M = Ti). An asterisk indicates a spinning sideband or surface methoxide. The appearance of a distinct resonance 7' is indicative of phenoxide coordination to Ti. The inset highlights the downfield shift of resonance 4 in 3 relative to 2 due to methyl ether coordination to Ti.](image-url)
Material 3 can be expected to be an active and selective epoxidation catalyst as a result of its rigid immobilization within a pseudotetrahedral geometry, which is often cited as a requirement for heterogeneous epoxidation catalysis on Ti centers.17,18,33–35 Epoxidation rates and selectivities for several unfunctionalized alkenes (Table 2) were measured at 60 °C using CHP or TBHP as oxidants and dry octane as solvent. Material 3 catalyzes cyclohexene epoxidation with both TBHP and CHP; rates were slightly higher with CHP. Cyclohexene epoxidation on catalyst 3 typically proceeded to >95% conversion and >95% selectivity to the epoxide (based on hydroperoxide). Epoxidation of 1-octene on 3 and cyclohexene on 4b occurs at a very low rate, and as such, unproductive thermal and surface-catalyzed decomposition of the hydroperoxide accounts for much of the total conversion, and the selectivity decreases to 30% and 66%, respectively.

### Table 2. Representative Epoxidation Results at 60 °C in Octane

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**Literature Comparisons**

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$^a$ General reaction conditions are given in the Experimental Section. $^b$ $k$ determined from $r = k[\text{cat}][\text{P}][\text{A}]$ based on the observed epoxide production rate. TOF$_0 = \left(\text{moles of epoxide/mole of Ti}/\text{hour}\right)$ extrapolated to initial time. $^c$ Total conversion of peroxide at a given time. $^d$ A 5 μmol sample of 4b, 5 mL of octane, and 0.65 mmol of cyclohexene. $^e$ Only 7 mL of octane used to increase all reactant concentrations. $^f$ Reaction was extremely rapid, and insufficient data could be collected to calculate a meaningful rate constant.
Regardless, for all surface coverages and substrates, no side products (i.e., other C$_6$ oxides for cyclohexene) were detected by gas chromatography, essentially ruling out contributions from competing radical pathways. Also, epoxidation of cis-stilbene proceeded with complete retention of configuration to only cis-stilbene oxide, which confirms that a nonradical mechanism is responsible for epoxidation in our catalysts.

The epoxidation of cyclooctene and 1-octene proceeded at the expected rates relative to those of cyclohexene. The observed relative substrate reactivity was cyclooctene > cyclohexene > 1-octene, consistent with attack of the alkene at an electrophilic coordinated peroxide oxygen. The successful epoxidation of stilbene and cyclooctene indicates the absence of significant steric constraints at the active site. Table 2 also shows that cyclohexene epoxidation rates on catalyst 3 resemble or exceed those reported on state-of-the-art Ti-based heterogeneous epoxidation catalysts. Of particular note is the high TOF reported for entry 20. This material is a highly dispersed grafted species that provides good active site accessibility. However, as for other catalysts in Table 2, the TOF of the catalyst in entry 21 decreases as the surface density increases, suggesting that site isolation may be incomplete.

**Kinetic Analysis.** Figure 3A shows catalytic turnovers to epoxide (TON = moles of epoxide/mole of Ti) in CHP and excess cyclohexene as a function of reaction time using catalyst at 60 °C. Figure 3B shows epoxide yields based on peroxide (X = moles of epoxide/initial mole of peroxide). Neither plot shows an induction period, indicating that the reaction conditions are much more dilute than most in the literature using Ti-based catalysts; thus, rates reported here on material underestimate those expected at the higher reactant concentrations of comparative studies. We note that our reaction conditions are much more dilute than most in the literature using Ti-based catalysts; thus, rates reported here on material underestimate those expected at the higher reactant concentrations of comparative studies.

Figure 3. Cyclohexene epoxidation in octane at 60 °C catalyzed by 3: [Figure 3A](#) 3-203, (●) 3-146, (●) 3-139, (◨) 3-75, (◨) 3-51, (▲) 3-25. Open symbols are for CHP, and closed symbols are for TBHP. (A) TON to epoxide, CHP oxidant. Lines are fit using the first-order rate law proposed in the text. (B) Conversion X to epoxide, CHP oxidant. Lines are obtained by linear regression of the data points shown. The slope is proportional to the pseudo-first-order rate constant $k_1$. (C) Second-order rate constants $k_2$ for CHP (---) and TBHP (●) plotted against total Ti concentration in the reactor. Lines are obtained by linear regression of the data points shown. The slope represents the intrinsic rate constant for epoxidation of cyclohexene with each oxidant.

The data in Figure 3B show a linear dependence of \( \log(1 - X) \) on time at all peroxide conversions, and the pseudo-first-order rate constants, proportional to the slope of these lines, are unaffected by increasing reaction time. These results indicate that catalyst deactivation, leaching, or inhibition by products do not occur. Consistent with this linear behavior, the deliberate addition of an excess of coproduct alcohol at initial time did not influence reaction rates or reactant conversions.

The lack of kinetic inhibition by alcohol products suggests a strong preference for binding peroxide reactants instead of abundant alcohol products on Ti active centers. First-order behavior was observed previously on Ti−silsesquioxane complexes, and early Ti−SiO₂ catalysts, but many initially active epoxidation catalysts deactivate with reaction time, an effect that has been ascribed to inhibition by alcohol products.

Grafted calixarene−Ti IV catalysts reported here can be stored in ambient air without detectable changes in epoxidation rates and selectivities compared to freshly prepared materials. This stability is in marked contrast to the sensitivity of conventional Ti epoxidation catalysts to atmospheric moisture during storage, which causes irreversible restructuring around Ti centers. As with other epoxidation catalysts stable during storage, such as grafted Ti(OMe)₅, traces of water during catalysis led to lower selectivities and rate constants with increasing reaction time, apparently as a result of acid-catalyzed epoxide hydrolysis, which forms chelating diols that bind strongly to Ti centers.

This situation is avoided by conducting the reaction in the presence of molecular sieves, as we have performed here.

**Experimental Evidence for Single-Site Behavior.** Silica gel surfaces are known to exhibit a distribution of acid sites, and this can influence the number and type of ligands at a grafted metal center. In contrast, we expect the bonding around the Ti center in 3 to be largely controlled by the multidentate calixarene ligand, which should force the Ti into a pseudotetrahedral coordination geometry, as in the single-crystal X-ray diffraction structure of precursor 4b. To test this hypothesis, we examined the epoxidation of cyclohexene using 3 as catalyst, with a wide range of surface concentrations of Ti active sites from 0.03 to 0.25 nm⁻².

Catalytic activity in a single-site material depends only on the total concentration of Ti centers and not their surface concentration. In Figure 3A, turnover frequencies for cyclohexene epoxidation (TOF = TON/time) extrapolated to initial times show identical TOFs (slope) on catalysts with surface densities between 0.06 and 0.25 nm⁻². This behavior is usually considered sufficient to prove single-site behavior, but here we compare rate constants over the entire time course of reaction to unequivocally prove the single-site behavior of 3 as catalyst.

In view of the linearity of the conversion data when plotted as \( \log(1 - X) \) and the large excess of cyclohexene reactants used, we use a rate law given by \( r = k_2[x][P][A] \), where \( [P] \) is the concentration of hydroperoxide (mol/L) and \( [A] \) is the concentration of alkene (mol/L), which is essentially unchanged during reaction. The first-order dependence on hydroperoxide was established above, and the dependence on alkene was verified independently. At the reaction conditions given in the Supporting Information, the initial TOF increased linearly from 34 to 65 to 160 h⁻¹ as the cyclohexene concentration was raised from 18 to 48 to 138 mM (Supporting Information Figure S4). The final concentration is approximately that of the reaction system employed in this paper. The catalyst concentration [cat] is expressed as millimoles of Ti per reactor volume (L). The rate was obtained from the integrated form of the rate expression \( \ln(1 - X) = k_2[x][P][A] = k_2[A] = k_2t \). A graph of \( \ln(1 - X)[A]^{-1} \) versus time has a slope of \( k_2 \) and a zero intercept. A plot of \( k_2 \) as a function of [cat] (Figure 3C) leads to a straight line through the origin for single-site materials and a curve of decaying slope for conventional grafted Ti catalysts.

The value of \( k_2 \) was determined for each catalyst using this methodology, and the linear dependence shown by the data in Figure 3C unequivocally shows that intrinsic rate constants depend only on the number of Ti centers in the reactor and not on the calixarene−Ti IV surface coverage, an accepted hallmark of single-site catalysts. Also, the use of intrinsic rate constants instead of initial TOF ensures that these materials behave identically at all conversions, rather than only during initial contact with reactants. The rate constants \( k \) in Figure 3C are 11.1 ± 0.3 M⁻²s⁻¹ for TBHP and 25 ± 2 M⁻²s⁻¹ for CHP over the entire range of calixarene−Ti IV surface coverages (0.03−0.25 nm⁻²).

This demonstrated single-site behavior differs markedly from that reported on other Ti-based catalysts, for which epoxidation turnover rates (per Ti center) either decrease monotonically with increasing Ti concentration or sharply decrease above a certain value, apparently because metal centers increasingly reside at inactive or inaccessible surface sites or because oligomers form with less active and selective Ti−O−Ti connectivity. We propose that the steric bulk and geometric constraints provided by the calixarene ligand prevent metal center clustering or nonuniform coordination geometries even near saturation coverage.

**Identification of Catalytically Relevant Species.** Control experiments first sought to confirm the heterogeneous nature of the catalysis observed and to determine the stability of 3 against leaching during reaction. The homogeneous precursor 4b (Table 2, entry 1) and its tert-butoxy analogue showed >20-fold lower activity than the grafted material 3. The addition of pure silica to a reaction mixture containing homogeneous material 4b synthesizes 3 in situ, as determined by TGA and UV−vis spectroscopy of recovered solids, and leads to a concomitant increase in epoxidation rates (Supporting Information Figure S5a). This illustrates the importance of Ti−O−Si connectivity for epoxidation catalysis and demonstrates synergy between grafted precursors and the support.

In several experiments, a reaction mixture of catalyst 3−138, CHP, and cyclohexene was filtered at 60 °C after reaction for 10 min (TON ≈ 30). The filtrate showed no detectable epoxidation activity over 1 h (Figure S5b). Also, the addition of pure silica to the filtrate, a procedure that leads to in situ immobilization of 3 and to the appearance of catalytic activity


when homogeneous 4b is present in solution, did not lead to detectable epoxidation products after an additional 1 h at 60 °C.

The absence of leaching cannot be demonstrated unequivocally from the observed noncatalytic character of the filtrate because of the low epoxidation rates on homogeneous catalyst 4b and on its tert-butoxy analogue. Recycle experiments (Figure S5c), in which the catalyst is washed with hot anhydrous octane, dried, weighed, and used for another catalytic cycle, showed very similar reaction rates and conversions in subsequent catalytic runs (within 5%), confirming the absence of permanent deactivation or leaching of sites. Also, addition of a second aliquot of hydroperoxide after 100% conversion (Figure S5d) led to an identical conversion rate, confirming that the catalyst remains active in the presence of all reaction products and is not, perhaps, reactivated by the recycling process.

Analysis of the solids recovered after filtration also did not detect elemental or structural changes indicative of leaching of calixarenes or of Ti-containing species. TGA and Ti ICP-MS independently did not detect changes in the carbon or titanium content within their respective accuracies (±5%). UV-vis spectroscopy (Figure 2A) and 13C CP/MAS NMR (Supporting Information Figure S6) of the solids recovered after catalysis showed that 3 was essentially unaltered during reaction. When catalysis was deliberately performed under deactivating conditions (without molecular sieves), the loss in activity was paralleled by a loss in calixarene content by both UV-vis and 13C CP/MAS NMR experiment, aromatic calixarene resonances were replaced by alkoxide resonances. These data, coupled with an unchanged Ti content by ICP-MS, suggest that diole formation from epoxide hydrolysis leads to the displacement of the calixarene ligand with concomitant loss of catalytic activity.

Deliberate calixarene removal by combustion in dry air at 500 °C produces catalyst 6-146, which shows behavior essentially similar to that of the grafted and calcined titanocene catalyst reported by Maschmeyer et al.34 Under identical conditions,

the initial epoxide TOF on 6-146 was 504 h⁻¹ as compared to 416 h⁻¹ on 3-146, but unlike the grafted calixarene catalyst, epoxide production essentially stopped at an 80% yield of epoxide. After this point, epoxide consumption continued and a small fraction of the epoxide was decomposed (likely to ring-opened product). At identical conditions, catalyst 3 continued to >98% conversion with 100% selectivity. The relative increase in initial TOF for 6-146 presumably reflects the removal of steric limitations around the Ti center imposed by the calixarene ligand. At high conversions, the absence of a calixarene ligand results in strong inhibition by coproduct alcohol, as reported previously.34

Taken together, these results indicate that material 3 is stable with respect to ligand loss and does not leach into solution as less active homogeneous species, such as titanium alkoxides, which can potentially form during reaction. We conclude, therefore, that the immobilization of active species 3 is permanent under the conditions reported here and that it is required for efficient and selective epoxidation catalysis.

Proposed Reaction Intermediate. We propose here structure 7 for the active species as consistent with catalytic and characterization data, but we note that the bonding mode of the Ti-peroxide intermediate in heterogeneous epoxidation catalysts remains unclear.48–51

The bound peroxide may be hydrogen-bonded with adjacent calixarene phenols or surface silanols as proposed for epoxidation in the presence of polar protic species,50 or the intermediate could be an η² complex as proposed for the active site in grafted complexes51 and observed in model compounds.52 Also, while we cannot rule out insertion of the peroxide into the Ti=O–Si bond,4,18,49 we believe that insertion into one of the Ti–OC bonds is more likely. Ti–OC bonds have been previously shown to be more kinetically labile than Ti–Osi bonds in the presence of TBHP.53 Particular to our system, the three-fold multiplicity of Ti–OC bonds relative to a single Ti–Osi bond favors the proposed structure 7, and cleavage of a Ti–OC bond will be further favored by release of conformational strain in the metallocalixarene. This strain is apparent in the elliptical conformation of the calixarene macrocycle in the structure of 4b as determined by single-crystal X-ray diffraction.11 Finally, 7 maintains the Ti–Osi connection required for the observed heterogeneous catalysis.

Many factors, including steric constraints at the metal,18,33,53 diffusion to the active site,20,39,54 and support hydrophobicity,36,42,55 influence epoxidation catalysis on Ti centers, but alkene epoxidation rates are expected to increase with increasing electron-withdrawing power of oxo ligands.40 In this context, the main role of the calixarene ligand in 3 in epoxidation catalysis is to withdraw electron density from the peroxide oxygens in a reaction intermediate such as 7, rendering such oxygens more electrophilic and prone to alkene attack.

We suggest that calixarene ligands in 3 perform two simultaneous functions by (i) introducing steric constraints that prevent dimerization and formation of weaker Ti–O–Ti Lewis acids18,34,56 and (ii) promoting Lewis acidity at the Ti center
relative to conventional alkoxide or Cp ligands. This stronger Lewis acidity reflects the effective delocalization of electron density away from the Ti center via Ti–O–Ph π-bonding; this mechanism is analogous to that invoked for the increased Lewis acidity in early-transition-metal complexes with siloxy ligands relative to those with alkoxy ligands. A similar argument has also been invoked for the high activity and stability of grafted transition-metal alkyls as polymerization catalysts.

Conclusions

We demonstrate here a novel route for the synthesis of isolated Ti–oxo complexes via grafted metallocalixarenes and their use as highly active and selective epoxidation catalysts. The single-site nature of the observed electronic transitions and catalysis on these materials confirms our premise that the steric bulk of calixarene ligands prevents oligomerization events leading to inactive and unselective Ti–O–Ti structures. The coordination environment enforced by the cone calixarene ligand leads to strong Lewis acidity and to faster epoxidation rates compared with those in conventional Ti centers with alkoxy and Cp ligands. The calixarene ligand provides steric constraints around each Ti center, which leads to remarkable stability during ambient storage and epoxidation catalysis, as well as to 100% selectivity to epoxide products. We expect this to be a general method for creating surface organometallic species since we have shown that the calixarene–TiIV connectivity has been maintained from the crystallographically characterized precursor to the active catalytic species. We are currently exploiting the diversity and flexibility made available by calixarene ligands and the possibility of modifying the support properties for additional improvements in epoxidation rates and selectivities on grafted metallocalixarene materials.

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Supporting Information Available: Catalysis method and substrate purification details, characteristic TGA of 3-146 (Figure S1), characteristic nitrogen physisorption data (Figure S2), characteristic fluorescence emission spectra (Figure S3), investigation of reaction order in alkene concentration (Figure S4), silica addition (Figure S5a), filtration (Figure S5b), recycle (Figure S5c), and second aliquot (Figure S5d) catalysis experiments, and UV–vis (Figure S6a) and 13C NMR (Figure S6b) comparison for fresh, active, and deactivated catalysts. This material is available free of charge via the Internet at pub.acs.org.

Supporting Information Available: