Terminal Titanyl Complexes of Tri- and Tetrametaphosphate: Synthesis, Structures, and Reactivity with Hydrogen Peroxide
Julia M. Stauber, and Christopher C. Cummins*
Department of Chemistry, Massachusetts Institute of Technology, 77 Massachusetts Avenue, Cambridge, Massachusetts 02139

Received February 1, 2017; E-mail: ccummins@mit.edu

ABSTRACT: The synthesis and characterization of tri- and tetrametaphosphate titanium(IV) oxo and peroxy complexes is described. Addition of 0.5 equiv [OTi(acac)2]2− to dihydrogen tetrametaphosphate ([P4O12H2]2−) and monohydrogen trimetaphosphate ([P3O9H]2−) provided a bis(μ2-κ2,κ3) tetrametaphosphate titanyl dimer, [OTiP4O12]2+ (1, 70% yield), and a trimetaphosphate titanyl acetylacetonate complex, [OTiP3O9(acac)]2− (2, 59% yield). Both 1 and 2 have been structurally characterized, crystallizing in the triclinic P1 and monoclinic P21 space groups, respectively. These complexes contain Ti=O units with distances of 1.624(7) and 1.644(2) Å, respectively, and represent rare examples of structurally characterized terminal titanyls within an all-oxygen coordination environment. Complexes 1 and 2 react with hydrogen peroxide to produce the corresponding peroxotitanium(IV) metaphosphate complexes, [O2TiP4O12]2+ (3, 61% yield), and [O2Ti2P3O9(acac)]2− (4, 65% yield), respectively. Both 3 and 4 have been characterized by single crystal X-ray diffraction studies, and their solid-state structures are presented. Complex 3 functions as an oxygen atom transfer (OAT) reagent capable of oxidizing phosphorus(III) compounds (P(OMe)3, PPh3) and SMet2 at ambient temperature to result in the corresponding organic oxide with regeneration of dimer 1.

■ INTRODUCTION

Oxidative degradation of organic ligands is a challenge faced in many synthetic systems as most oxidation catalysts require ligand environments capable of supporting reactive high-valent metal oxos. Furthermore, the deactivation of high-valent metal oxos through the formation of unreactive dimeric or oligomeric μ-oxo species is an additional common challenge. In heterogeneous systems, these issues are circumvented through site isolation of discrete metal oxos within inorganic matrices that function as thermodynamically stable ligand environments. An example of one such system is Enichem’s highly efficient TS-1 oxidation catalyst, a titanium based microporous molecular sieve in which Ti(IV) ions are isolated within an entirely silicate based framework. The success of TS-1 and other industrial oxidation catalysts has driven synthetic chemists to study molecular models of heterogeneous systems. Kortz et al. have studied titanium(IV) oxo and hydroxo substituted polyoxometalates (POMs) as models of Ti single site catalysts due to the structural analogy of POMs and metal oxide surfaces. These oxidatively stable titanium(IV) POM complexes have served as molecular models for studying mechanisms of selective oxidation. With the present work, we sought to investigate the suitability of similarly all-inorganic metaphosphate rings (cyclic oligomers of the PO3 anion) for stabilization of the titanyl functional group in an all-oxygen coordination environment.

■ RESULTS AND DISCUSSION

Synthesis of Titanyl Complexes. The [PPN]+ salts of both dihydrogen tetrametaphosphate ([PPN]2[P4O12H2]) and monohydrogen trimetaphosphate ([PPN]2[P3O9H]) can be prepared in multigram quantities under open air conditions by protonation of their corresponding metaphosphate salts with strong acid. We have previously shown that the [P4O12H2]2− acid can be used to synthesize metal complexes through protolysis of basic ligands such as [N(SiMe3)2]2−, leading to the facile preparation of tetrametaphosphate tin(II) ([SnP4O12])2− and chromium(II) ([CrP4O12]2−) salts. Employing the strategy of protolytic ligand replacement, we chose to investigate the reactivity of both the [P4O12H2]2− and [P3O9H]2− acids with titanyl bis(acetylacetonate) ([OTi(acac)2])13. Even though its structure is dimeric, [OTi(acac)2]2− provides a commercially available source of the titanyl (OTi) unit that bears basic ligands potentially susceptible to protolytic replacement by the metaphosphate acids. Acetylacetonate, the byproduct formed through protonation, is easily separable from the desired metaphosphate salts, making the preparation of titanyl metaphosphate complexes convenient and straightforward.

Scheme 1. Synthetic protocol for the preparation of [PPN]4[OTiP4O12]2+ ([PPN]4[I]).

Treatment of 1 equiv [PPN]2[P4O12H2] with 0.5 equiv [OTi(acac)2]2− in an acetone/MeCN mixture at 23 °C afforded a new species, [OTiP4O12]2+ (1), after 12 h of reaction time (Scheme 1). Analytically pure X-ray quality crystals of [PPN]4[I] were easily isolated from the acetylacetonate byproduct in 70% yield after concentration of the crude reaction mixture and storage at 23 °C overnight. The identity of 1 was established through an X-ray diffraction study, and the solid-state structure is shown in Figure S58 (left). The 31P{1H} NMR spectrum of dimer 1 features a singlet located at δ = −27.55 ppm (Figure S1), which is shifted slightly upfield from that of [P4O12H2]2−. The Ti=O stretching frequency observed by FTIR at 969 cm−1 (Figure S5) is in line with data reported for other terminal titanyl complexes reported in the literature.14−17 and falls within the characteristic range (972−890 cm−1) associated with the Ti=O oscillator.18 The formula of the [OTiP4O12]2+ anion was also confirmed by ESI-MS(−) with a m/z value of 189.88 (Figure S7, calc’d: 189.89). [PPN]4[I] crystallized in the triclinic space group, P1̅1, with two [PPN]+ cations and half of the [OTiP4O12]2+ dimer in the asym-
metric unit. As shown in Figure 1, the complex adopts a [MP_4O_12]_2 coordination geometry in which two tetrametaphosphate rings form a cofacial sandwich around the pair of titanyl ions resembling the structure of [CP_4TiP_4O_12]_2- reported by Ishii et al. We reported previously that the chromium(II) complex of tetrametaphosphate forms just such a dimeric cofacial sandwich. In I, the titanium centers reside 0.559 Å above the equatorial four-membered oxygen plane of the tetrametaphosphate such that two κ^2-[P_2O_12]_2^4- ligands bridge each Ti center to approximate D_2h point group symmetry. Dimer I contains a Ti-Ti distance of 4.233(9) Å, with a titanyl Ti=O distance of 1.624(7) Å. While structurally characterized anionic titanium(IV) oxo complexes are rare, the observed Ti=O distance in I agrees well with corresponding metrical parameters reported for other neutral titanium(IV) oxo complexes with distances spanning 1.61 to 1.68 Å. The Ti=O distance observed in I is consistent with triple bond character as predicted by the titanium and oxygen triple bond covalent radii, as well as the electronic considerations associated with d^0 transition metal oxos within tetragonal ligand fields.

**Scheme 2.** Synthetic protocol for the preparation of [PPN][OTiP_3O_9(acac)] ([PPN][2]).

As in the case of the dihydrogen tetrametaphosphate, the [PPN][2]OTiP_3O_9(acac) acid reacted with 0.5 equiv [OTi(acac)]_2 in an acetone/MeCN mixture under open air conditions to generate a titanyl metaphosphate complex, [PPN][2]OTiP_3O_9(acac)] ([PPN][2], Scheme 2, Figure 2), with loss of one equiv acetylacetone. Complex 2 was isolated in 59% yield by crystallization through concentration of the crude reaction mixture and allowing the resulting solution to stand at 23 °C. After 12 h, large colorless crystals formed that were isolated and washed with acetone to afford analytically pure [PPN][2].

The 31P{^1H} NMR spectrum of 2, recorded in acetonitrile, features a singlet located at δ = −18.43 ppm that is assigned to the three phosphorus atoms of the [P_2O_10]^{12-} ligand (Figure S8). The presence of a single broad signal can be ascribed to rapid rotation of the [P_2O_10]^{12-} unit on the NMR time scale in solution, and is therefore represented as a weighted average. Collection of the 31P{^1H} NMR spectrum at −35 °C resolved the broad singlet into a doublet and a triplet with an AX_2 splitting pattern located at δ = −19.13 (^2J_PP = 17.2 Hz) and −18.20 ppm (^2J_PP = 17.2 Hz) in a 2:1 integral ratio (Figure S29). The doublet corresponds to the phosphorus atoms flanking the two equatorially bound oxygen atoms, and the triplet is assigned to the phosphorus bound to the axially coordinated oxygen atom of the [P_3O_9]^{12-} ligand.

![Figure 1](image1.jpg)

**Figure 1.** Solid-state structure of [OTiP_2O_9(acac)]_2+ (1) [OTiP_3O_9(acac)]^{2-} rendered with PLATON\textsuperscript{25} with thermal ellipsoids at the 50% probability level, and with [PPN]^+ cations and solvent molecules of crystallization omitted for clarity. Selected interatomic distances (Å) and angles (°): Ti1-O1 1.652(15), Ti1-O2 1.999(12), Ti1-O3 1.9939(13), Ti1-O4a 1.9844(13), Ti-O5a 2.0014(13), Ti1-Ti1a 4.233(9), Σ(Oeq-Ti-Oeq) 341.8(9).

![Figure 2](image2.jpg)

**Figure 2.** Solid-state structure of [OTiP_2O_9(acac)]^{2-} (2) rendered with PLATON\textsuperscript{25} with thermal ellipsoids at the 50% probability level, and with [PPN]^+ cations and solvent molecules of crystallization omitted for clarity. Selected interatomic distances (Å) and angles (°): Ti1-O1 1.644(2), Ti1-O2 2.033(2), Ti1-O3 2.049(18), Ti1-O4 2.0145(19), Ti1-O5 2.0113(19), Ti1-O6 2.3211(19), Z(Oeq-Ti-Oeq) 355.08(2).

An X-ray diffraction study confirmed the solid-state structure of 2 as a terminal titanyl complex with one [P_2O_9]^{12-} and one acetylacetone ligand bound to the titanium(IV) center to complete its oxygen based octahedral coordination environment. Figure 2 displays a thermal ellipsoid plot of the anion from the [PPN][2] salt. As a result of the strong trans influence imparted by the oxo ligand,\textsuperscript{20} the Ti1–O5 distance of 2.3211(19) Å is significantly longer than expected for a Ti–O single bond (1.99 Å) as predicted by the titanium and oxygen single bond covalent radii,\textsuperscript{23} suggesting this interaction is at best a weak one and that O5 may be simply in proximity of the Ti center by virtue of being part of the trimetaphosphate ring. Consistent with multiple bonding to a terminal oxo ligand, the IR spectrum of 2 displays the Ti=O oscillator at ν = 953 cm\textsuperscript{−1} (Figure S11). This stretching frequency is 16 cm\textsuperscript{−1} lower in energy than that of 1, and consistent with the slightly longer Ti=O distance of 2 (1.644(2) Å) compared with that of 1 (1.6252(13) Å). In contrast to 2, the position trans to the oxo ligand is vacant in dimer 1, and thus 1 should be able to maximize the Ti=O multiple bond in accord with both its shorter bond distance and higher energy stretching frequency. The formula of 2 was also confirmed by ESI-MS(−) with a m/z value of 300.91 (Figure S14, calc’d: 300.82), which corresponds to the monoanionic complex, [OTiP_3O_9]^{-}, with loss of the acac\textsuperscript{−} ligand under ESI-MS(−) conditions.

Dimer 1 was prepared under open atmosphere conditions and is air stable both in solution and as a solid. Dimer 1, however, reacted with excess water (ca. 50–100 eq) in acetonitrile at 23 °C over the course of 5 min to form a new species exhibiting a higher order splitting pattern in its 31P{^1H} NMR spectrum consistent with an

![Figure 3](image3.jpg)

**Figure 3.** $^{17}$O NMR spectrum of 2 after treatment with excess (40 eq) $^{17}$OH$_2$ (MeCN, 68 MHz, 20 °C, externally referenced to D$_2$O). Inset: ESI-MS(−) of 2 after treatment with excess $^{17}$OH$_2$ (MeCN, 3200 V).
The water stability of (ca. 200 equiv) over the course of 24 h in acetonitrile (Figure S30). When an acetonitrile solution containing this species was heated to 80 °C under vacuum, the $^{31}$P{[^1]H} NMR spectrum of the resulting residue shows resonances corresponding to dimer 1, revealing the reversibility of the aquation reaction (Figure S43). When a solution of 1 with excess water was allowed to stand at 23 °C for 12 h, complete conversion of the initial species to a new product was observed (Figure S40).

To gain more insight into the identity of these products, a reaction was carried out using $^{17}$O enriched water. The $^{17}$O NMR spectrum of an acetonitrile solution of 1 treated with excess water (ca. 40 eq) $^{17}$OH$_2$ displays signals located at $\delta$ 1060 and 930 ppm (Figure S41), which fall within the range expected for terminal titanium(IV) oxo (1000–1100 ppm) $^{15,28,29}$ and hydroxo ligands (700–1000 ppm), $^{29}$ respectively, and are in line with our assignment of this species as the mixed O=Ti/Me$_2$O$_2$ bistratetrametaphosphate complex. When a sample was permitted to age for 12 h, these resonances were observed to decrease in intensity with new signals appearing at $\delta$ 758 and 566 ppm (Figure S42). These new signals are found within the region expected for Ti(IV) $\mu^2$-O (650–850 ppm) and $\mu^3$-O bridged species (450–600 ppm), $^{30}$ suggesting that after an extended period of time in the presence of a large excess of H$_2$O, dimer 1 is converted to $\mu$-oxo bridged oligomers.

Complex 2 exhibits similar behavior to dimer 1 in being air stable in both the solid state, and in solution. In contrast to 1, however, 2 is stable to excess water as judged by the unchanged $^{31}$P NMR spectrum when a sample was left to stand in the presence of water (ca. 200 equiv) over the course of 24 h in acetonitrile (Figure S30). The water stability of 2 afforded the opportunity to study the titanyl oxygen exchange with $^{17}$OH$_2$ and $^{18}$OH$_2$ by ESI-MS(−) and $^{17}$O NMR spectroscopy. Comba and Merbach have reported that the oxygen exchange rate for terminal oxo ligands in systems of hydrolyzed Ti(IV) is extremely fast as measured by $^{17}$O NMR fast-injection and line-broadening techniques. $^{29}$ Consistent with these studies, the titanyl oxygen of complex 2 exchanged readily with both $^{16}$OH$_2$ and $^{18}$OH$_2$ as evaluated by ESI-MS(−) studies. As a result of this facile exchange, a $^{17}$O NMR spectrum of 2 (Figure 3) was obtained after spiking a saturated acetonitrile solution of the compound with $^{17}$OH$_2$ (ca. 40 eq). The solution $^{17}$O NMR spectrum displays one sharp signal located at $\delta$ 1086 ppm that is assigned to the $\gamma$-oxo of 2, and consistent with $^{17}$O NMR studies of related terminal titanyl complexes. $^{15,28}$

The formation of 1 and 2 from [OTi(acac)$_2$]$_2$ is noteworthy in illustrating that the tri- and tetrametaphosphates are capable of breaking up the titanium(IV) bis-$\mu$-oxo core of [OTi(acac)$_2$]$_2$ to result in the formation of terminal Ti=O units. Titanium(IV) oxo species are typically susceptible to hydrolysis in the presence of moist air as they have the propensity to dimerize and oligomerize forming unreactive species containing Ti=O-Ti linkages, $^{3,26}$ so the formation of a well-defined, air-stable terminal titanyl complex is unusual. Structurally characterized examples of complexes containing terminal titanyl units are known and typically contain nitrogen based macrocyclic ligands $^{31,32}$ such as porphyrins, $^{16}$ phthalocyanines, $^{33}$ and tacn derivatives (tacn = 1,4,7-triazacyclononane), $^{17,34}$ however, examples containing an oxygen-only coordination environment at titanium are extremely rare. $^{35}$ Such examples include [OTi(OSMe$_2$)$_2$]$^{12+}$ $^{3,36}$ and [OTi(CO)$_3$]$^{1-}$ $^{3,37}$ complexes, and a sandwich-type 8-tungsto-2-arsenate(III) POM system ([OTi$_2$(α-$\alpha$-$\alpha$-As$_2$W$_6$O$_{24}$)$_2$]$^{12+}$) $^{3,38}$ Additionally, direct evidence for the formation of terminal [O=Ti(acac)]$^{12+}$ units in dilute acidic aqueous solutions of Ti(IV) has been observed. $^{39}$ The IR spectrum of such a solution contains a band at 975 cm$^{-1}$ attributed to the Ti=O vibration, while the $^{17}$O NMR spectrum features a resonance located at $\delta$ 1028 ppm that is assigned to the $\gamma$-Oxygen. These data are in line with the IR and $^{17}$O NMR spectroscopic signatures of complexes 1 and 2, suggesting that the ligand field exerted by metaphosphates may be similar to that of water.

**Reactivity of Titanyl Complexes with Hydrogen Peroxide.**

The conversion of titanium(IV) oxo to titanium(IV) $\eta^2$-peroxo complexes through reaction with H$_2$O$_2$ is a well understood transformation $^{38}$ that has been thoroughly studied in both molecular $^{39,40}$ and heterogeneous $^{41}$ systems. This reactivity is of particular interest as hydrogen peroxide is used in many titanium silicate systems in catalytic oxidation reactions of organic substrates. $^{6,41-43}$ This transformation is not only relevant to oxidation processes, but because these reactions are typically accompanied by the appearance of an intense orange color, titanyl reactivity with hydrogen peroxide has been the basis for a colorimetric H$_2$O$_2$ sensing technology. $^{44}$

Treatment of dimer 1 with 2 equiv urea-H$_2$O$_2$ (UHP) or Me$_3$SiOOSiMe$_3$ in acetonitrile resulted in a color change from colorless to orange, concomitant with the clean formation of the corresponding tetrametaphosphate peroxotitanium(IV) dimer ([O$_2$Ti$_2$P$_2$O$_8$])$_2$$^{4-}$, 3) with loss of 2 equiv H$_2$O or Me$_3$SiOOSiMe$_3$, respectively. When UHP was used as the peroxide source (Scheme 3), the reaction was complete within 10 min, while the corresponding reaction with Me$_3$SiOOSiMe$_3$ was much more sluggish and required 12 h for completion. Both reactions were monitored with $^{31}$P NMR spectroscopy wherein progress was associated with growth in of a singlet at $\delta$ $-$28.13 ppm (Figure S15). Additionally, the reaction was followed by UV-Vis spectroscopy by noting the appearance and growth of the absorption located at $\lambda_{max}$ = 379 nm (Figure S20). The product was isolated (61% yield) as orange blocks obtained by layering diethyl ether onto an acetonitrile solution of the complex.

An X-ray diffraction study confirmed the identity of this product as a peroxotitanium(IV) $k^2$-tetrametaphosphate dimer [PPN][O$_2$Ti$_2$P$_2$O$_8$]$_2$ (3, with the peroxo ligands bound $\eta^2$ to each titanium center. Figure 4 (left) shows a thermal ellipsoid plot of the anion from this salt. The O10-O20 peroxo ligand lies in the O4-Ti1b-O2a plane. The identity of 3 was also confirmed by ESI-MS(−), with a m/z value of 197.95 (Figure S21, calc’d, 197.89). The Ti=O

![Scheme 3. Treatment of [OTiP$_2$O$_4$]$^{3-}$ (1) with 2 equiv H$_2$O$_2$ to yield [O$_2$Ti$_2$P$_2$O$_8$]$^{4-}$ (3) with loss of 2 equiv H$_2$O.](image-url)
Fig. 4. Solid-state structure of [O₂TiP₂O₇]^{4-} (3) rendered with PLATON visualized with PLATON. (1) with thermal ellipsoids at the 50% probability level, and with [PPN⁺] cations and solvent molecules of crystallization omitted for clarity. Selected interatomic distances (Å) and angles (°): Ti1-O1 1.815(8), Ti1-O20 1.795(8), Ti1-O1 1.896(13), Ti1-O2 1.935(12), Ti1-O3b 1.986(13), Ti1-O4a 1.963(12), O20-O10 1.486(6), Ti1-Ti1b 4.23(1), Σ(0eq,Ti-Oeq) 352.0(7).

vibration at 969 cm⁻¹ in the IR spectrum of 1 is absent in the spectrum of 3, and a new stretch attributed to the peroxo O–O vibration is located at 856 cm⁻¹ (Figure S18), and is consistent with a side-on π coordination mode. The strong absorption at λ_max = 379 nm in the UV-Vis spectrum of 3 is assigned to the p₉(O2)→dₓ(Ti) ligand to metal charge-transfer (LMCT) transition, and compares well with other LMCT bands that have been observed for similar peroxotitanium(IV) complexes. Like 1, dimer 3 is air stable in both the solid state and in solution.

When the reaction of 1 with UHP was monitored by 31P NMR spectroscopy at 5 °C in acetonitrile, an intermediate evincing an A₂B₂C₂' splitting pattern (Figure S35) was observed before complete conversion to 3 occurred. This intermediate is assigned as the Os/Ti/Ti(O–O) bistetrametaphosphate complex, and completely converts to complex 3 after 80 min at 5 °C, and can only be observed at low temperature (≤ 5 °C).

Scheme 4. Reaction of [OTiP₂O₇(acac)]^{2-} (2) with H₂O₂ to yield [O₂TiP₂O₇(acac)]^{2-} (4) with loss of H₂O.

Complex 2 also reacted cleanly with H₂O₂(aq) or Me₃SiO(OH)SiMe₃ to generate a peroxotitanium(IV) trimetaphosphate acetylacetonate complex, [PPN]₂[O₂TiP₂O₇(acac)] (Scheme 4, Figure 5), which was isolated in 65% yield after crystallization. 4 is characterized by a sharp singlet in its 31P{¹H} NMR spectrum (Figure S22), a strong stretch at νₚ₂ₒ = 903 cm⁻¹ in its IR spectrum (Figure S25), and an absorption located at λ_max = 403 nm (Figure S27, ε = 674 M⁻¹cm⁻¹) in the visible region of the electromagnetic spectrum. Crystals of [PPN]₂[4] suitable for an X-ray diffraction study were grown as orange blocks obtained by layering Et₂O onto an acetone/tritluoromethane solution of the complex. [PPN]₂[4] crystallized in the monoclinic space group P2₁, and Figure 5 shows a thermal ellipsoid plot of the anion that displays the peroxo ligand in an π⁻ binding mode in which the O–O moiety is coincident with the O2–Ti1–O3 plane of the [P₂O₇]^{2-} and acac⁻ ligands. The structure determination revealed unremarkable Ti–Oₐ₇peroxo and O–Oₐ₇distances of 1.850(4), 1.825(3), and 1.443(6) Å, respectively.

Examples of molecular peroxotitanium(IV) complexes that serve as oxidants towards organic substrates are rare due to the well-documented stability of the titanium-peroxo structural unit, and therefore its corresponding lack of oxidizing capacity. It is instead believed that titanium(IV) hydroperoxo species play the active role in substrate oxidation within heterogeneous systems, and therefore the preparation of well-defined titanium hydroperoxo complexes has been targeted by synthetic chemists.

We were unable to observe a stable titanium(IV) hydroperoxo complex, and 1 and 2 did not show any signs of catalytic activity toward organic substrates in the presence of H₂O₂ (See Section 13 in the Supporting Information for experimental details). Both results are likely due to the rapid conversion of 1 and 2 to 3 and 4, respectively, upon addition of H₂O₂ without a long-lived hydroperoxo intermediate for interception by a substrate. No reaction was observed when 3 and 4 were independently treated with substrates relevant to TS-1 oxidation processes such as diphenylmethanol, phenol, and cyclooctene (See Section 14 in the Supporting Information for experimental details). These results are consistent with the lack of catalytic activity observed for 1 and 2 in the presence of H₂O₂.

Scheme 5. Oxygen atom transfer (OAT) reactions from [O₂TiP₂O₇(acac)]^{2-} (3) to P(OMe)₃, PPh₃, and SMe₂ to generate the corresponding organic oxide and [OTiP₂O₇]^{2-} (4) under the following conditions: a) 23 °C, 10 min, MeCN; b) 23 °C, 20 min, MeCN; c) 23 °C, 2 h.

We did, however, test the ability of 3 and 4 to behave as stoichiometric oxidants towards phosphorus(III) compounds and sulfides. Treatment of 3 with P(OMe)₃, PPh₃, and SMe₂ at ambient temperature resulted in the formation of the corresponding organic oxide as well as clean regeneration of 1 as determined by 31P and ¹H NMR analyses (Scheme 5). Treatment of 4 with PPh₃, however, required heating at 70 °C to generate OPPH₃ and 3. Related vanadium(V) peroxo systems are thermodynamically powerful but kinetically sluggish OAT reagents, and in this respect they are reminiscent of nitrous oxide.

■ CONCLUSIONS

The present metaphosphate complexes represent rare examples of air stable Ti(IV) oxo/peroxo systems containing entirely oxygen based coordination environments at titanium. The metaphosphate ligands support the formation of terminal titanyls as opposed to the...
more commonly encountered $\mu$-oxo bridged oligomeric structures, making complexes 1–4 molecular models of site isolated titanyl and peroxo units found within heterogeneous oxidation catalysts.

**Experimental Section**

**General Methods.** Unless otherwise stated, all manipulations were performed under open air conditions in the fume hood. [PPN][2[P2O5]2H2] \(^{11}\) and [PPN][2[P2O5]2H] \(^{12}\) were prepared as previously reported. All reagents were purchased from Sigma Aldrich (urea, H2O2, [OTi(acac)]\(_2\), 30% H2O2), or Gelest (Me2SiOSiMe3) and used as received. Acetone (H2O content < 0.5 w/w%) was purchased from Macron Fine Chemicals, and acetonitrile was purchased from Acros Organics and both were used without further purification. 17O[H2O] (97% enriched) and 18OH2 (97% enriched) were purchased from Cambridge Isotope Laboratories, and used as received. IR spectra were recorded on a Bruker Tensor 37 Fourier transform IR (FTIR) spectrometer. Deuterated solvents (D2O, CD3CN) were purchased from Cambridge Isotope Laboratories, and used as received. NMR spectra were obtained on a Varian Inova-500 NMR spectrometer with an Oxford Instruments Ltd. superconducting magnet, Bruker Avance 400 instruments equipped with Magnex Scientific superconducting magnets, or on a Varian Mercury 300 spectrometer with an Oxford Instruments Ltd. superconducting magnet. 1H and 13C\(^{1}H\) NMR spectra are referenced to residual protio-solvent signals. 31P\(^{1}H\) NMR chemical shifts are reported with respect to an external reference (85% H3PO4, 0.0 ppm), and 17O NMR chemical shifts are externally referenced to D2O (0.0 ppm). NMR spectra were simulated using the program gNMR version 5.1. ESI-MS(−) data were recorded on a Waters Q-TOF micro mass spectrometer using a source temperature of 100 °C and a desolvation temperature of 150 °C. Mass spectrometry samples were run in neat acetonitrile at concentrations < 1 μM, and the data were processed using the program mMass Version 5.4.1. UV-Vis spectra were collected on an Ocean Optics USB4000 spectrophotometer with a DT-Mini-2GS UV-vis-NIR light source. Elemental analyses were performed by Robertson Microlit Laboratories, Inc (http://www.robertsonmicrolit.com). X-ray crystallographic details are provided in Section 15.1 of the Supporting Information.

[PPN]\(_{4}[OTiP_{2}O_{4}]_{2}\) [][PPN]\(_{4}[2][1]\). To a stirring suspension of [PPN][2[P2O5]2H2] (350 mg, 0.251 mmol, 1.00 eq) in acetone (7 mL) was added [OTi(acac)]\(_2\) (72 mg, 0.14 mmol, 0.55 eq) as a solid. MeCN (10 mL) was added dropwise until a homogeneous solution was obtained. The pale yellow reaction mixture was allowed to stir at room temperature (23 °C) for a total of 12 h, at which point the mixture was filtered through a pad of Celite to remove unreacted [OTi(acac)]\(_2\). The filtrate was concentrated to ca. 0.5 mL under reduced pressure, and dried under reduced pressure to afford analytically pure [OTiP\(_2\)O\(_4\)]\(_{2}\) (310 mg, 0.210 mmol, 92%). X-ray diffraction quality crystals of [PPN][2[OTiP\(_2\)O\(_4\)] were grown by concentration of [PPN][2[OTiP\(_2\)O\(_4\)] to 0.5 mL, and the filtrate was allowed to stand at 23 °C for 12 h, during which time large colorless crystals formed that were isolated and washed with acetone to afford analytically pure [PPN][2]. Overall yield of crystals: 199 mg, 0.134 mmol, 59%. The solid-state structure determination revealed the presence of one MeCN molecule of crystallization per [PPN][2[OTiP\(_2\)O\(_4\)] complex. Elem. Anal. (%) found (calc’d) for C\(_{79}H_{116}N_{10}O_{26}P_{4}Ti_{2}\) [([PPN][2[OTiP\(_2\)O\(_4\)]MeCN)]: C, 59.42 (56.00); H, 3.41 (3.38); N, 15.87 (15.74); P, 5.27 (5.18); Ti, 11.42 (11.50). ESI-MS(−) of [OTiP\(_2\)O\(_4\)]\(_{2}\) as a cream colored powder (310 mg, 0.210 mmol, 92%). X-ray diffraction quality crystals of [PPN][2[OTiP\(_2\)O\(_4\)] were grown by concentration of the crude reaction mixture by ca. 90% and allowing the resulting mixture to stand at 23 °C. After 12 h, large colorless crystals formed that were isolated and washed with MeCN to afford analytically pure [PPN][2]. Overall yield of crystals: 199 mg, 0.134 mmol, 59%. The solid-state structure determination revealed the presence of one MeCN molecule of crystallization per [PPN][2[OTiP\(_2\)O\(_4\)] complex. Elem. Anal. (%) found (calc’d) for C\(_{79}H_{116}N_{10}O_{26}P_{4}Ti_{2}\) [([PPN][2[OTiP\(_2\)O\(_4\)]MeCN)]: C, 59.42 (56.00); H, 3.41 (3.38); N, 15.87 (15.74); P, 5.27 (5.18); Ti, 11.42 (11.50). ESI-MS(−) of [OTiP\(_2\)O\(_4\)]\(_{2}\) on [PPN][2]. To a stirring suspension of [PPN][2[P2O5]2H] (300 mg, 0.228 mmol, 1.00 eq) in acetone (7 mL) was added [OTi(acac)]\(_2\) (66 mg, 0.13 mmol, 0.55 eq) as a solid. MeCN (10 mL) was added dropwise until the reaction mixture became homogeneous. The pale yellow reaction mixture was allowed to stir at ambient temperature (23 °C) for a total of 6 h, at which point the pale yellow solution was filtered through a pad of Celite to remove unreacted [OTi(acac)]\(_2\). All volatiles were then removed under reduced pressure affording a pale yellow residue. The yellow residue was triturated with Et2O (10 mL), and the Et2O was subsequently decanted from the resulting solids. The resulting solids were suspended in acetone (5 mL), and filtered away from the pale yellow supernatant. The solids were washed with acetone (3 mL) and dried under reduced pressure, affording [PPN][2[OTiP\(_2\)O\(_4\)] as a colorless powder (70% enriched) and 18OH2 (10%), and the Et2O was subsequently decanted from the resulting solids. The resulting solids were suspended in acetone (5 mL), and filtered away from the pale yellow supernatant. The solids were washed with acetone (3 mL) and dried under reduced pressure, affording [PPN][2] as a cream colored powder (310 mg, 0.210 mmol, 92%). X-ray diffraction quality crystals of [PPN][2[OTiP\(_2\)O\(_4\)] were grown by concentration of the crude reaction mixture by ca. 90% and allowing the resulting mixture to stand at 23 °C. After 12 h, large colorless crystals formed that were isolated and washed with acetone to afford analytically pure [PPN][2]. Overall yield of crystals: 199 mg, 0.134 mmol, 59%. The solid-state structure determination revealed the presence of one MeCN molecule of crystallization per [PPN][2[OTiP\(_2\)O\(_4\)] complex. Elem. Anal. (%) found (calc’d) for C\(_{79}H_{116}N_{10}O_{26}P_{4}Ti_{2}\) [([PPN][2[OTiP\(_2\)O\(_4\)]MeCN)]: C, 62.28 (62.61); H, 4.76 (4.65); N, 2.72 (2.77). ESI-MS(−) of [OTiP\(_2\)O\(_4\)]\(_{2}\) [([PPN][2[OTiP\(_2\)O\(_4\)]MeCN)]: C, 62.28 (62.61); H, 4.76 (4.65); N, 2.72 (2.77). ESI-MS(−) of [OTiP\(_2\)O\(_4\)]\(_{2}\) [([PPN][2[OTiP\(_2\)O\(_4\)]MeCN)]: C, 62.28 (62.61); H, 4.76 (4.65); N, 2.72 (2.77). ESI-MS(−) of [OTiP\(_2\)O\(_4\)]\(_{2}\) [([PPN][2[OTiP\(_2\)O\(_4\)]MeCN)]: C, 62.28 (62.61); H, 4.76 (4.65); N, 2.72 (2.77).
To the concentrated solution was added Et₂O (15 mL) in order to precipitate orange solids. The solids were isolated by filtration, washed with Et₂O (15 mL), and dried under reduced pressure affording [PPN]₄[3] as an orange solid. Yield: 85 mg, 0.029 mmol, 86%.  

[PPN]₂[O₂TiP₃O₈(acac)] ([PPN]₂[4]). To a stirring colorless solution of [PPN][2] (100 mg, 0.0677 mmol, 1.00 eq) was added a solution of 30% H₂O₂ (10 µL, 0.098 mmol, 1.4 eq) via 10 µL microsyringe. The color of the solution immediately changed from colorless to bright orange upon complete addition. The orange solution was allowed to stir at ambient temperature for a total of 10 min, at which point all volatiles were removed under reduced pressure. The resulting orange residue was triturated with THF (5 mL), and then the THF was subsequently decanted away from the orange solids. The solids were then dissolved in MeCN (1 mL), and then Et₂O (3 mL) was added until the solution became cloudy. The orange solution was then filtered through a piece of microfiber filter paper and allowed to stand at ambient temperature (23 °C) for 12 h. The pale orange mother liquor was removed and the crystals were dried under reduced pressure to afford analytically pure [PPN]₂[4] as an orange solid. Yield: 74 mg, 0.0340 mmol, 50%.  

IR (ATR, cm⁻¹): ν(C=O): 1719.00 (s, C=O(acac)): 1719.00 (s, C=O(acac)). IR (ATR, cm⁻¹): ν(Ti=O): 900.00 (s, Ti=O(acac)): 900.00 (s, Ti=O(acac)). MS (–) conditions, and only the [O₂TiP₃O₈(acac)] complex including the acac ligand is not detectable under ESI–MS(–) conditions.  

Acknowledgement This work was supported by NSF under the NSF Center CHE-1305124. Yanfeng Jiang is thanked for assistance with X-ray crystallography, and Wesley Transue and Shiyu Zhang are thanked for assistance with VT-NMR experiments.

Supporting Information Available: Experimental details and characterization data for all compounds including crystallographic information for [PPN][1], [PPN][2], [PPN][3], and [PPN][4] are provided in the Supporting Information. This material is available free of charge via the Internet at http://pubs.acs.org.

References


(17) Bodmer, A.; Jeske, P.; Weyhermueller, T.; Wieghardt, K.; Dubler, E.; Schmelke, H.; Niiber, M. Mono- and Dinuclear Titanium(III)/Titanium(IV)
Complexes with 1,4,7-trimethyl-1,4,7-triazaacyclononane (L). Crystal Structures of Compositionally Diverse Green and a Blue Form of [Li[TiCl3][O][N(CS)2]2]: [Li[Ti(OH)2]BeCl2] and [Li2Ti2O3F2(f)2][PF6]. Inorg. Chem. 1992, 31, 3737–3748.


A new pair of air stable terminal titanyl complexes supported by tri- and tetrametaphosphate ligands is reported. These compounds react with H$_2$O$_2$ to generate the corresponding peroxo complexes with loss of water, and represent rare examples of terminal titanium(IV) oxos stabilized by an all-oxygen ligand field.