

Cite this: *Chem. Sci.*, 2020, 11, 10729

All publication charges for this article have been paid for by the Royal Society of Chemistry

Redox chemistry and H-atom abstraction reactivity of a terminal zirconium(IV) oxo compound mediated by an appended cobalt(I) center†

Hongtu Zhang,^a Gillian P. Hatzis,^a Diane A. Dickie,^b Curtis E. Moore^a and Christine M. Thomas^{*,a}

The reactivity of the terminal zirconium(IV) oxo complex, $\text{O}\equiv\text{Zr}(\text{MesNP}^i\text{Pr}_2)_3\text{CoCN}^t\text{Bu}$ (**2**), is explored, revealing unique redox activity imparted by the pendent redox active cobalt(I) center. Oxo complex **2** can be chemically reduced using Na/Hg or $\text{Ph}_3\text{C}^\bullet$ to afford the $\text{Zr}^{\text{IV}}/\text{Co}^0$ complexes $[(\mu\text{-Na})\text{OZr}(\text{MesNP}^i\text{Pr}_2)_3\text{CoCN}^t\text{Bu}]_2$ (**3**) and $\text{Ph}_3\text{COZr}(\text{MesNP}^i\text{Pr}_2)_3\text{CoCN}^t\text{Bu}$ (**4**), respectively. Based on the cyclic voltammogram of **2**, $\text{Ph}_3\text{C}^\bullet$ should not be sufficiently reducing to achieve the chemical reduction of **2**, but sufficient driving force for the reaction is provided by the nucleophilicity of the terminal oxo fragment and its affinity to bind Ph_3C^+ . Accordingly, **2** reacts readily with $[\text{Ph}_3\text{C}][\text{BPh}_4]$ and Ph_3CCl to afford $[\text{Ph}_3\text{COZr}(\text{MesNP}^i\text{Pr}_2)_3\text{CoCN}^t\text{Bu}][\text{BPh}_4]$ (**5**) and $\text{Ph}_3\text{COZr}(\text{MesNP}^i\text{Pr}_2)_3\text{CoCl}$ (**6**), respectively. The chemical oxidation of **2** is also investigated, revealing that oxidation of **2** is accompanied by immediate hydrogen atom abstraction to afford the hydroxide complex $[\text{HOZr}(\text{MesNP}^i\text{Pr}_2)_3\text{CoCN}^t\text{Bu}]^+$ (**9**)[†]. Thus it is posited that the transient $[\text{OZr}(\text{MesNP}^i\text{Pr}_2)_3\text{CoCN}^t\text{Bu}]^+ [\text{2}]^+$ cation generated upon oxidation combines the basicity of a nucleophilic early metal oxo fragment with the oxidizing power of the appended cobalt center to facilitate H-atom abstraction.

Received 2nd August 2020
Accepted 4th September 2020

DOI: 10.1039/d0sc04229c

rsc.li/chemical-science

Introduction

Metal oxo compounds are proposed to play key roles in many essential chemical transformations such as oxidation reactions,^{1–8} oxygen transfer reactions,^{9,10} C–H activations,¹¹ and oxygen evolution reactions (OER).¹² In order to understand these important oxygenation/oxidation reactions, great efforts have been made to isolate and study the electronic structure and reactivity of metal oxo compounds.

Despite the high oxophilicity¹³ and low d electron counts that should favor metal-oxo multiple bond formation, examples of terminal oxo complexes of group 4 metals (Ti, Zr, Hf) are relatively scarce, especially for Zr and Hf. Instead, group 4 metals have a strong tendency to dimerize and form oxo-bridged clusters with bridging oxides.^{14–16} The “hard” Lewis acid nature of group 4 metals enhances the basicity of their oxo ligands, which therefore exhibit a strong preference to bridge two or more metal centers.¹⁷ As a strategy to synthesize terminal

oxo compounds with group 4 metals, sterically bulky ligands have been shown to effectively inhibit dimerization through the oxo ligand. In the limited examples of Ti oxo compounds, macrocyclic tetraaza ligand frameworks^{18–20} and coordinatively saturated octahedral geometries^{21,22} have been adopted to stabilize reactive Ti oxo moieties. Ti or Zr oxo species in (pseudo)tetrahedral geometries are often isolated in anionic form and stabilized by alkali cations,^{23–25} a strategy that also works for macrocyclic terminal oxo species.²⁶ A notable example of a neutral titanocene terminal oxo complex, which was quite unstable and readily decomposed to bridging oxo species, was reported by Andersen and coworkers.²⁷

Compared with Ti terminal oxo compounds, there are even fewer Zr terminal oxo species reported. Bergman and coworkers conducted detailed mechanistic and reactivity studies of a transient zirconocene terminal oxo species, generated *in situ* through heating the Zr hydroxide precursor $\text{Cp}^*\text{ZrPh}(\text{OH})$ ($\text{Cp}^* = \eta^5\text{-C}_5\text{Me}_5$).^{30,31} Subsequently, Parkin isolated the first neutral Zr terminal oxo compound, $(\eta^5\text{-C}_5\text{Me}_4\text{Et})_2\text{Zr}(\text{O})(\text{NC}_5\text{H}_5)$ (Fig. 1), generated by $2e^-$ oxidation of Zr^{II} precursor $\text{Cp}^{*f}\text{Zr}(\text{CO})_2$ with N_2O ($\text{Cp}^{*f} = \eta^5\text{-C}_5\text{Me}_4\text{Et}$).^{29,32} This zirconocene oxo compound showed rich nucleophilic reactivity across a wide range of substrates such as water, iodomethane, silanes, and ketones. To our knowledge, Parkin's zirconocene oxo compound was the only example of a structurally characterized neutral Zr terminal oxo compound prior to our recent report of a Zr/Co hetero-bimetallic oxo compound (Fig. 1, *vide infra*).²⁸

^aDepartment of Chemistry and Biochemistry, The Ohio State University, 100 W. 18th Ave, Columbus, OH 43210, USA. E-mail: thomasc@chemistry.ohio-state.edu

^bDepartment of Chemistry, University of Virginia, 409 McCormick Road, PO Box 400319, Charlottesville, VA 22904, USA

† Electronic supplementary information (ESI) available: Experimental procedures and characterization data, spectroscopic and cyclic voltammetry data, additional computational and crystallographic details. CCDC 2020655–2020661. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/d0sc04229c



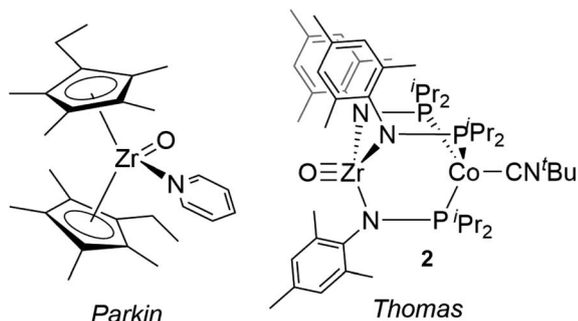


Fig. 1 Terminal Zr oxo complexes reported by Parkin (left)²⁹ and Thomas (right).²⁸

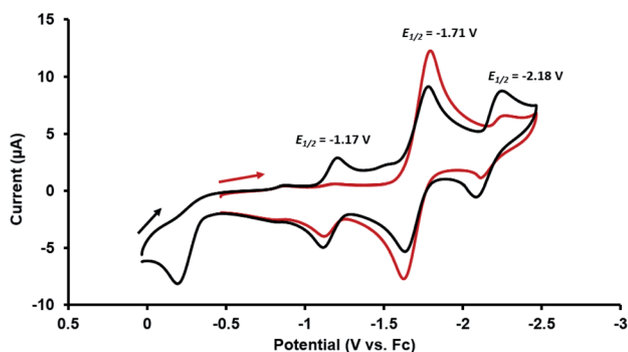
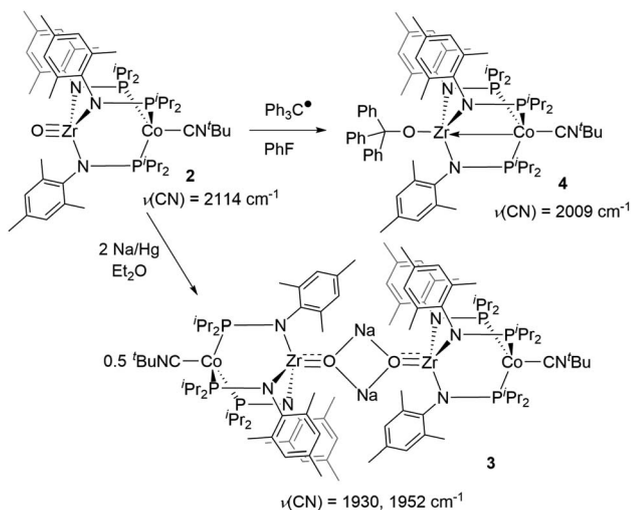


Fig. 2 Cyclic voltammograms of **2** (in 0.3 M [^tBu₄N][PF₆] in THF; scan rate: 100 mV s⁻¹) scanning cathodically starting from the open circuit potential (red) and starting from 0 V (black) vs. Fc/Fc⁺.



Scheme 1 Chemical reduction of **2** using Na/Hg and Ph₃C[•].

The synthesis of a bimetallic terminal Zr oxo compound adds to the growing body of literature on bimetallic complexes, which have received interest for applications in small molecule activation and catalysis.^{33–36} Recently, our group reported the early/late heterobimetallic compound (THF)Zr(MesNPⁱPr₂)₃CoCN^tBu (**1**),³⁷ derived from (THF)Zr(MesNPⁱPr₂)₃CoN₂ via

replacement of N₂ with CN^tBu. The tightly bound CN^tBu apical ligand sterically protects the Co site and considerably alters the reaction profile of the bimetallic compound. In studies of the reactivity of (THF)Zr(MesNPⁱPr₂)₃CoN₂, many bond activation processes involve simultaneous substrate binding to both metal centers facilitated by dissociation of the labile N₂ ligand,^{39–44} however, the isoelectronic Zr^{IV}/Co^{-I} complex **1** lacks an accessible coordination site on Co, requiring Zr to mediate substrate activation while the Co center acts as a redox active metal-ligand providing electrons for substrate reduction. This redox-active metalloligand approach was first showcased in oxidation reactions with organic azides to afford terminal Zr imido species.³⁷ More recently, complex **1** was shown to activate O₂ to afford the two-electron oxidized peroxy species (O₂)Zr(MesNPⁱPr₂)₃CoCN^tBu and to react readily with O-atom transfer reagents to afford the terminal oxo compound O≡Zr(MesNPⁱPr₂)₃CoCN^tBu (**2**), shown in Fig. 1.²⁸ With the appended Co center, complex **2** has the potential for further redox chemistry in addition to nucleophilic reactivity, a marked difference compared to the reaction space available to the monometallic zirconocene terminal oxo compound. Herein, we describe the reactivity and redox chemistry of complex **2**.

Results and discussion

Cyclic voltammetry of O≡Zr(MesNPⁱPr₂)₃CoCN^tBu (**2**)

Given the presence of a redox active Co^I center in the heterobimetallic Zr^{IV}/Co^I complex **2**, the redox behavior of this compound was examined using cyclic voltammetry. The cyclic voltammogram (CV) of **2** showed both oxidative and reductive features. When scanned cathodically starting from the open circuit potential (Fig. 2, red), the CV showed a reversible reduction at -1.71 V (vs. the ferrocene/ferrocenium redox couple, Fc/Fc⁺) assigned as the Co^{I/0} redox couple. If the cathodic scan was, instead, started at a more positive potential (0 V vs. Fc/Fc⁺, Fig. 2, black), an irreversible oxidation process was observed at -0.21 V. After scanning through potentials sufficiently oxidizing to carry out this oxidation process, two additional reversible waves appeared at -1.17 V and -2.18 V, revealing that upon oxidation of **2**, a subsequent chemical step occurred to generate a new stable Zr/Co complex that can readily access two reversible redox processes. The rich redox behavior of oxo compound **2** enabled by the ancillary Co center was further explored through chemical oxidation and reduction reactions and characterization of the resulting products.

Reduction of O≡Zr(MesNPⁱPr₂)₃CoCN^tBu (**2**)

First, the chemical reduction of oxo complex **2** was investigated through treatment of **2** with Na/Hg amalgam. Treatment of **2** with excess Na/Hg in diethyl ether solution resulted in the formation of [(μ-Na)OZr(MesNPⁱPr₂)₃CoCN^tBu]₂ (**3**, Scheme 1). During the course of the reduction, the reaction mixture converted from a bright green suspension to a homogeneous orange solution, from which the light-yellow crystalline product **3** was isolated as the major product. The ¹H NMR spectrum of **3** showed 6 paramagnetically shifted resonances in a pattern



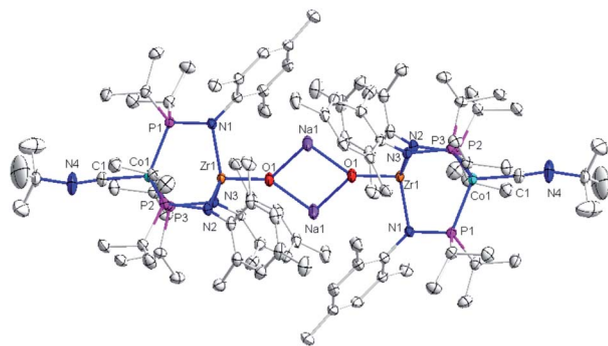


Fig. 3 Displacement ellipsoid (50%) representation of anionic oxo compounds **3**. Hydrogen atoms and solvate molecules were omitted for clarity. Selected interatomic distances (Å): Zr1–Co1, 2.8174(3); Zr1–O1, 1.8589(15).

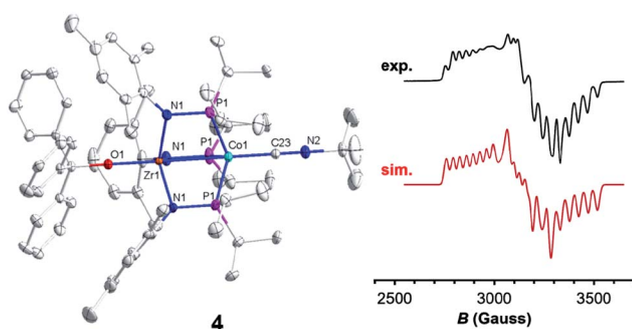
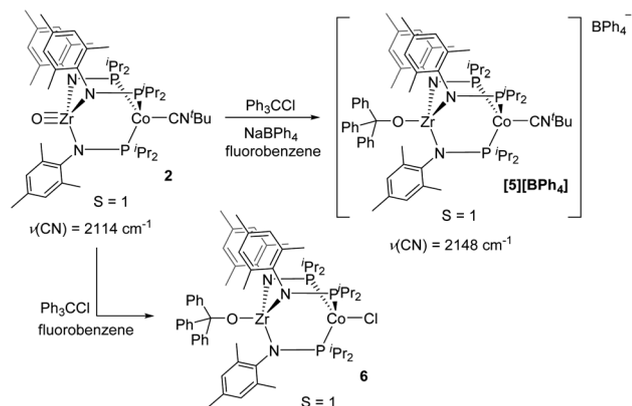


Fig. 4 Displacement ellipsoid (50%) representations (left) and X-band EPR spectra (right, experimental: black, top and simulated: red, bottom) of **4** in fluorobenzene at 30 K (power attenuation = 30 dB, modulation amplitude = 10 G, modulation frequency = 100 kHz). Hydrogen atoms and solvate molecules have been omitted for clarity. Selected interatomic distances (Å) of **4**: Zr1–Co1, 2.7223(5); Zr1–O1, 1.987(2). EPR simulation parameters: $g = 2.00, 2.11, \text{ and } 2.33$; $A = 130, 109, \text{ and } 82$ MHz.



Scheme 2 The reactivity of **2** with Ph_3C^+ reagents.

indicative of C_3 symmetry and similar to that of previously reported $S = 1/2$ tris(phosphinoamide) Zr/Co species.^{25,39,45} The solid-state IR spectrum of **3** has two $\text{C}\equiv\text{N}$ stretches at

1930 cm^{-1} and 1952 cm^{-1} . The $\text{C}\equiv\text{N}$ stretches are lower in frequency compared to the neutral oxo precursor **2** ($\nu(\text{CN}) = 2114 \text{ cm}^{-1}$)²⁸ as a result of stronger backbonding from the reduced Co center to the CN^tBu ligand.

The solid-state structure of compound **3** revealed a tetrametallic dimeric structure in which the Zr-bound anionic oxo ligands were bridged by two sodium cations to connect the two Zr/Co fragments (Fig. 3). Compound **3** bears structural and spectroscopic similarities to the previously reported carbonyl derivatives of the general form $[\text{OZr}(\text{MesNP}^i\text{Pr}_2)_3\text{CoCO}]^-$.²⁵ The Zr–Co distance in **3** is 2.8174(3) Å, approximately 0.2 Å shorter than that of the oxo compound **2** (3.0157(2) Å)²⁸ owing to both the more reduced Co center and the weakened Zr–O bonding due to the competing interaction between the oxo ligand and the alkali cation. The Zr–O bond distance in compound **3** is, indeed, elongated compared to the oxo compound (1.8589(15) Å in **3** vs. 1.7956(9) Å in **2**), indicative of disruption of the Zr–O multiple bonding. The Zr–O distance in **3** is comparable to that of $[(12\text{-crown-4})\text{Li}][\text{OZr}(\text{MesNP}^i\text{Pr}_2)_3\text{CoCO}]^-$,²⁵ which was assigned a $\text{Zr}\equiv\text{O}$ triple bond. Although one σ and two π bonds remain possible owing to the local C_3 symmetry of **3**, the interaction of the oxo ligand with the Na^+ cation as well as the trans influence of the appended Co center weakens the Zr–O bonding in **3** compared to **2**.

Triptyl radical also served as an effective inner sphere reductant for oxo compound **2**. A reaction between **2** and half an equivalent of Gomberg's dimer (3-triphenylmethyl-6-diphenylmethylidene-1,4-cyclohexadiene), which dissociates into Ph_3C^* in solution,^{46,47} afforded $\text{Ph}_3\text{COZr}(\text{MesNP}^i\text{Pr}_2)_3\text{CoCN}^t\text{Bu}$ (**4**) after stirring for two hours in fluorobenzene (Scheme 1). The solid-state structure of **4** confirmed addition of the triptyl radical to the Zr-bound oxo ligand to afford a new triphenyl methoxide functionality (Fig. 4). The paramagnetically shifted ^1H chemical shift pattern of **4** is similar to the isoelectronic $S = 1/2$ compound **3**, in support of an overall one-electron reduction process from **2** to **4**. The $\text{C}\equiv\text{N}$ stretch (2009 cm^{-1}) in the solid-state ATR IR spectrum of **4** is at higher frequency than isoelectronic compound **3**, which can be attributed to the difference in overall charge between the two $\text{Zr}^{\text{IV}}/\text{Co}^0$ molecules; however, the $\nu(\text{CN})$ of **4** remains significantly lower compared to $\text{Zr}^{\text{IV}}/\text{Co}^{\text{I}}$ oxo compound **2** (2114 cm^{-1}).²⁸

Since the poor solubility of compound **4** precluded measurement of its magnetic moment in solution, EPR spectroscopy was used to confirm its $S = 1/2$ spin state. The X-band EPR spectrum of **4** exhibited a rhombic signal with simulated g values of 2.00, 2.11, and 2.33 with an average $g = 2.15$ (Fig. 4), consistent with an $S = 1/2$ system. The eight-line hyperfine splitting pattern associated with each g tensor is attributed to coupling to the ^{59}Co nucleus ($I = 7/2, A = 130, 109, 82$ MHz), and therefore spin localization on the Co center. The combined spectral evidence confirms that the reduction of **2** is cobalt-centered with the unpaired electron in both **3** and **4** residing on the $d^9 \text{Co}^0$ center.

Although the potential of the $\text{Ph}_3\text{C}^*/\text{Ph}_3\text{C}^+$ redox couple suggests that Ph_3C^* is not a sufficiently strong reductant to reduce **2**,⁴⁸ the reaction is likely driven to completion by the strong affinity of the electrophilic Ph_3C^+ cation for the oxo



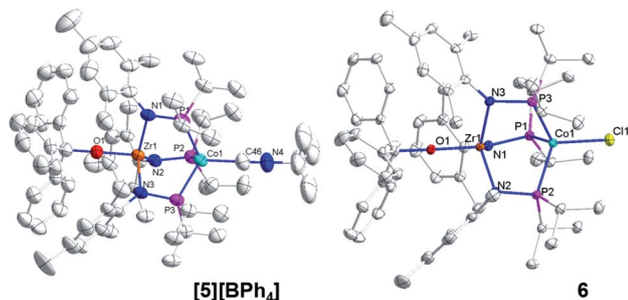
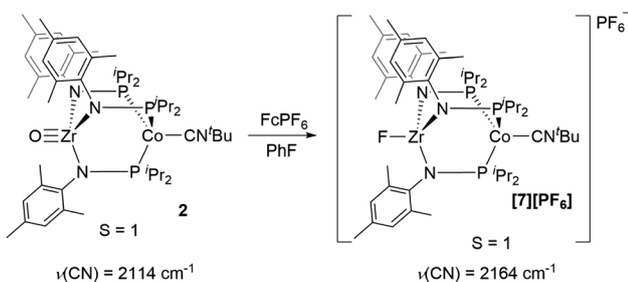


Fig. 5 Displacement ellipsoid (50%) representations of [5][BPh₄] and 6. Hydrogen atoms, solvate molecules, and the BPh₄[−] counteranion of [5][BPh₄] were omitted for clarity. Selected interatomic distances (Å): [5][BPh₄]: Zr1–Co1, 2.9212(4); Zr1–O1, 1.9301(15). 6: Zr1–Co1, 2.9005(3); Zr1–O1, 1.9558(8).



Scheme 3 Chemical oxidation of 2 with FcPF₆.

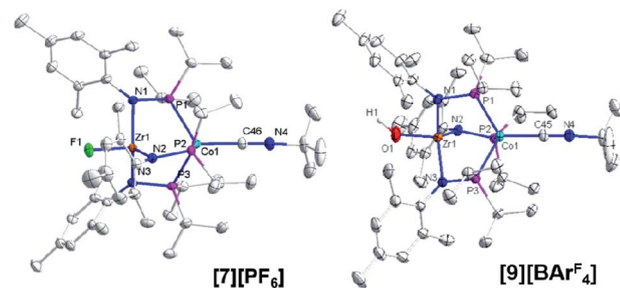


Fig. 6 Displacement ellipsoid (50%) representations of [7][PF₆] and [9][BARF₄]. Counteranions, solvate molecules, and hydrogen atoms except for the OH[−] group on [9][BARF₄] were omitted for clarity. Selected interatomic distances (Å): [7][PF₆]: Zr1–Co1, 2.7498(2); Zr1–F1, 1.9319(8). [9][BARF₄]: Zr1–Co1, 2.8396(3); Zr1–O1, 1.9209(13).

fragment. To test this hypothesis, complex 2 was treated directly with trityl cation reagents. The reaction between 2 and [Ph₃C][BPh₄] (generated *in situ* from NaBPh₄ and Ph₃CCl) afforded the bright green compound [Ph₃COZr(MesNPⁱPr₂)₃CoCN^tBu][BPh₄] ([5][BPh₄], Scheme 2). Compound [5][BPh₄] is the *S* = 1 (μ_{eff} = 2.95 B.M.), one-electron oxidized, isostructural analogue of 4 with BPh₄[−] as the counteranion. The paramagnetically shifted ¹H NMR chemical shift pattern of [5][BPh₄] is consistent with other C₃-symmetric Zr^{IV}Co^I tris(phosphinoamide) species such as oxo precursor 2.²⁸ The infrared C≡N stretching frequency of [5][BPh₄] (2148 cm^{−1}) is significantly higher than the Zr^{IV}/Co⁰

analogue 4 (2009 cm^{−1}) owing to the higher oxidation state of the Co center. The $\nu(\text{CN})$ is also considerably higher than that of isoelectronic 2 (2114 cm^{−1}), which can be attributed to the overall cationic charge of [5][BPh₄].

The addition of trityl cation to oxo compound 2 proceeds differently in the presence of a coordinating counteranion. A stoichiometric reaction between 2 and Ph₃CCl led to formation of the neutral blue product Ph₃COZr(MesNPⁱPr₂)₃CoCl (6, Scheme 2). Under these conditions, the tightly bound isocyanide ligand was displaced by the chloride ligand (Fig. 5). The solution magnetic moment of 6 (μ_{eff} = 2.81 B.M.) is consistent with an *S* = 1 species isoelectronic to compounds 2 and [5][BPh₄].

Compounds 4, [5][BPh₄], and 6 are structurally similar but with systematic changes in the intermetallic Zr–Co distances as a result of differences in the oxidation state of the cobalt center (Fig. 4 and 5). The Zr–Co distance (2.7223(5) Å) in the Zr^{IV}/Co⁰ compound 4 is the shortest in the 4–6 series, and is in good agreement with the Zr–Co distances reported for similar RO–Zr(MesNPⁱPr₂)₃Co–L species (R = H, alkyl, or aryl; L = CO, N₂; range = 2.6324(3)–2.7616(6) Å).^{25,39,45} The Zr–Co distances in Zr^{IV}/Co^I complexes [5][BPh₄] and 6 (2.9212(4) Å and 2.9005(3) Å, respectively) are elongated by approximately 0.2 Å compared to 4 as a result of diminished dative Co → Zr interactions from the less electron-rich oxidized Co^I center. The Zr–O distance in 4 (1.987(2) Å) is slightly elongated compared to [5][BPh₄] and 6 (1.9301(15) Å and 1.9558(8) Å, respectively), which may be reflective of a stronger *trans* influence from the more reduced Co center in 4.

Oxidation of O≡Zr(MesNPⁱPr₂)₃CoCN^tBu (2)

Next, we explored the chemical oxidation of 2, which we anticipated to be immediately followed by another chemical step based on the irreversibility observed in its CV. The chemical oxidation of 2 by FcPF₆ in fluorobenzene afforded the Zr-fluoride compound, [FZr(MesNPⁱPr₂)₃CoCN^tBu][PF₆] ([7][PF₆]), as the major product as a result of fluoride abstraction from the PF₆[−] anion (Scheme 3). [7][PF₆] is also generated when this reaction is performed THF, ruling out the fluorobenzene solvent as the source of the Zr-bound fluoride ion. The characterization data for [7][PF₆], including the $\nu(\text{CN})$ stretch (2164 cm^{−1}), the solution magnetic moment (μ_{eff} = 2.95 B.M.), and the range and pattern of chemical shifts in the ¹H NMR spectrum, allowed its formulation as a C₃-symmetric *S* = 1 Zr^{IV}Co^I species. While the signals for the PF₆[−] counterion were easily observed in the ³¹P NMR (−145.7 ppm, septet) and ¹⁹F NMR (74.5 ppm, doublet) spectra, the ¹⁹F NMR signal for the Zr–F ligand was not observed due to its proximity to the paramagnetic Co center.

Single crystal X-ray diffraction confirmed the connectivity of [7][PF₆] and revealed a Zr–Co interatomic distance of 2.7498(2) Å (Fig. 6), which is somewhat shorter than is typically observed for isoelectronic Zr^{IV}/Co^I compounds with oxygen-based ligands bound to Zr (*e.g.* Ph₃BOZr(MesNPⁱPr₂)₃CoCN^tBu, Zr–Co: 2.9793(6) Å).²⁸ This phenomenon is likely reflective of the more electronegative nature of the fluoride anion promoting a stronger Co → Zr dative interaction. On the other hand, the



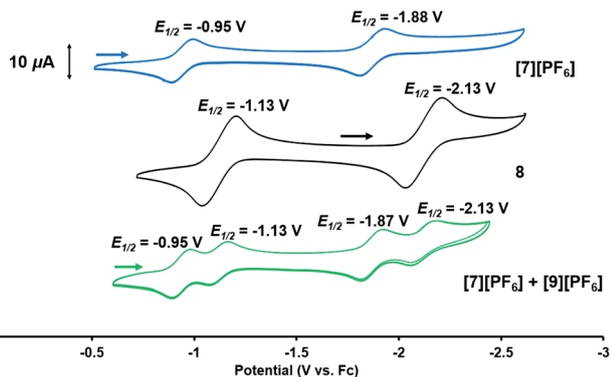
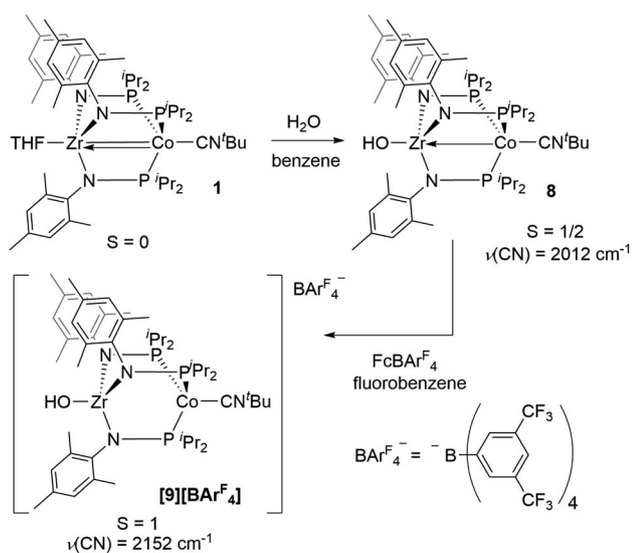


Fig. 7 Cyclic voltammograms (in 0.3 M [*t*Bu₄N][PF₆] in THF; scan rate: 100 mV s⁻¹) of compounds [7][PF₆] (blue), **8** (black), and a mixture of [7][PF₆] and [9][PF₆] obtained from the reaction of **8** with FcPF₆ (green).



Scheme 4 Synthesis of hydroxide compound **8** and its oxidation to form **9**.

Zr–Co distance 2.7498(2) Å is longer compared to that of the isoelectronic precursor ClZr(MesNPⁱPr₂)₃CoI (2.6280(5) Å),⁴⁹ likely due to the Co → CN^tBu back-donation to weaken dative Co → Zr interactions. Oxidation of **2** with ferrocenium salts containing less reactive tetraarylborate anions (*e.g.* FcBPh₄, FcBAr^F₄; BA^rF₄⁻ = tetrakis(3,5-bis(trifluoromethyl)phenyl) borate) led to mixtures of *S* = 1 products including both H-atom abstraction products ([9][BR₄], *vide infra*) and borane adducts R₃BOZr(MesNPⁱPr₂)₃CoCN^tBu²⁸ that were identified based on comparison of crude ¹H NMR spectra to previously characterized compounds.

Based on the formation of fluoride complex [7][PF₆] upon oxidation with FcPF₆, we initially hypothesized that the new reversible redox events that appeared at -1.17 V and -2.18 V in the CV of **2** after the irreversible oxidation step (Fig. 2) were attributed to rapid fluoride ion abstraction from the PF₆⁻ ions in the electrolyte solution. To probe this hypothesis, the CV of [7][PF₆] was collected (Fig. 7, blue). Two reversible cobalt-based

reduction events were observed in the CV of [7][PF₆] at -0.95 V and -1.88 V, which are significantly more positive and not consistent with the aforementioned features in the CV of **2**.

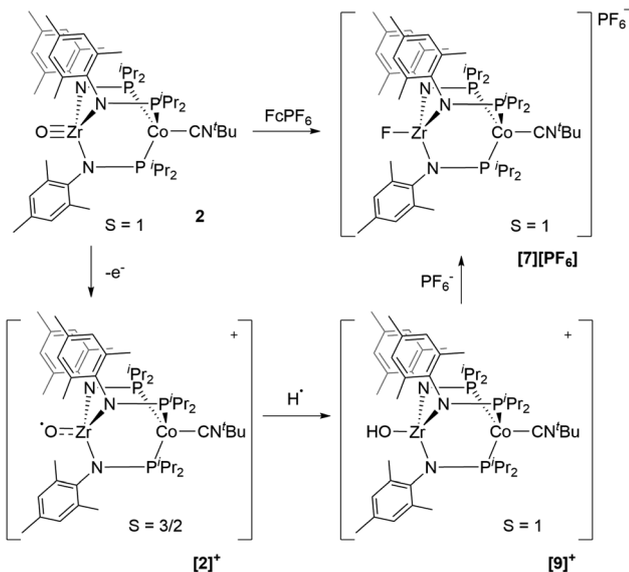
An alternative hypothesis to explain the new redox features in the CV of **2** and identify the species generated upon electrochemical oxidation of **2** was therefore warranted. We posited that upon oxidation, the newly generated cation, [OZr(MesNPⁱPr₂)₃CoCN^tBu]⁺ [**2**]⁺, might have both nucleophilicity and oxidizing power, making it prone to H-atom abstraction from the THF solvent to form a hydroxide complex of general form [HOZr(MesNPⁱPr₂)₃CoCN^tBu]⁺. To probe this hypothesis, the *S* = 1/2 hydroxide complex HOZr(MesNPⁱPr₂)₃CoCN^tBu (**8**) was synthesized *via* treatment of **1** with stoichiometric water in a reaction analogous to those previously reported with N₂- or CO-bound analogues of **1** (Scheme 4).^{25,45} The ¹H NMR spectrum of **8** contains eight broad paramagnetic resonances, but also features a diagnostic Zr–OH signal at -45.7 ppm comparable to resonances observed for the isoelectronic hydroxide compounds HOZr(MesNPⁱPr₂)₃CoN₂ (-52.8 ppm) and HOZr(MesNPⁱPr₂)₃CoCO (-40.6 ppm) (Fig. S21†).^{25,45} Single crystal X-ray diffraction confirmed the connectivity of **8**, including the hydroxide H atom, which was independently located in the difference map (Fig. S36†). The solid-state IR spectrum of **8** showed a ν(CN) at 2012 cm⁻¹ and a ν(OH) at 3701 cm⁻¹, and its solution magnetic moment was 1.64 B.M. Furthermore, the EPR spectrum of **8** features a rhombic signal centered at *g*_{av} = 2.14 (*g* = 2.00, 2.10, and 2.33) with a diagnostic eight line splitting pattern resulting from hyperfine coupling to the ⁵⁹Co nucleus (⁵⁹Co, *I* = 7/2, *A* = 132, 110, 100 MHz), further confirming the formulation of **8** as an *S* = 1/2 Zr^{IV}/Co⁰ complex (Fig. S23†).

The redox properties of hydroxide complex **8** were then investigated for comparison with the unknown species generated upon the oxidation of **2**. As shown in Fig. 7 and S24,† the CV of **8** reveals two reversible redox events, namely a Co⁰→I reduction at -2.13 V and a Co^I→0 oxidation at -1.13 V vs. Fc^{+/0}. These features are therefore quite similar to those observed following oxidation of oxo complex **2** (-1.17 V and -2.18 V), consistent with the hypothesis that H-atom abstraction is the rapid chemical step that follows oxidation.

One-electron oxidation of complex **8** with FcBA^rF₄ afforded the cationic *S* = 1 hydroxide compound [HOZr(MesNPⁱPr₂)₃CoCN^tBu][BA^rF₄]⁺ ([9][BA^rF₄]). In addition to the two signals for BA^rF₄⁻ anion, the ¹H NMR spectrum of [9][BA^rF₄] contains eight broad and paramagnetically shifted resonances, including a signal at -60.0 ppm corresponding to the Zr-bound hydroxide proton. The measured solution magnetic moment of [9][BA^rF₄] (*μ*_{eff} = 2.59 B.M.) is consistent with two unpaired electrons. The solid-state structure of [9][BA^rF₄] confirmed its connectivity and revealed an elongated Zr–Co interatomic distance compared to **8** (2.8396(3) Å vs. 2.6902(18) Å, Fig. 6). As expected, the redox features in the CV of cation [9]⁺ are superimposable with those of neutral complex **8** (Fig. S7†), but the open circuit potential of [9]⁺ is more positive (-0.87 V vs. -1.27 V vs. Fc^{+/0}).

Thus, we can definitively attribute the new signals observed in the CV of **2** following its oxidation to a cationic hydroxide species, [9]⁺, generated by an H-atom abstraction step by the





Scheme 5 Proposed transformation from oxo compound **2** to fluoride complex $[7][PF_6]$ based on experimental and computational study.

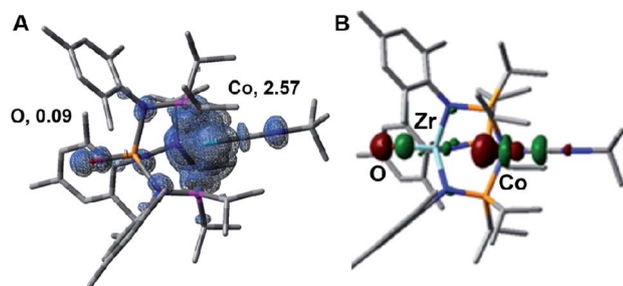


Fig. 8 (A) Calculated unpaired spin density surface of $[2]^+$ in the predicted quartet state, with Mulliken atomic spin densities on the cobalt and oxygen atoms indicated. (B) Calculated highest doubly occupied molecular orbital of **2**.

transiently generated cationic oxo species $[OZr(MesNP^iPr_2)_3-CoCN^tBu]^+$ ($[2]^+$). The proclivity of this oxidized compound to abstract H-atoms is unique to this heterobimetallic species, as a monometallic Zr oxo complex would not be sufficiently oxidizing to participate in such a proton-coupled electron transfer step. A remaining question, however, is why a cationic hydroxide complex such as $[9]^+$ is not isolated upon oxidation of **2** with $FcPF_6$ or, conversely, why the fluoride cation $[7]^+$ is not generated under electrochemical oxidation conditions. Indeed, the cleavage of a strong Zr–O bond to form fluoride complex $[7]^+$ was initially quite surprising. In a control reaction, no reaction was observed between oxo complex **2** and $[^tBu_4N][PF_6]$, revealing that fluoride ions are not abstracted from PF_6^- under non-oxidative conditions. Furthermore, solvent effects could be ruled out as oxidation of **2** with $FcPF_6$ in THF rather than fluorene did not lead to detectable formation of $[9]^+$. We therefore propose that one-electron oxidation to generate $[2]^+$ is immediately followed by rapid H-atom abstraction to form $[9]^+$

under both chemical and electrochemical oxidation conditions (Scheme 5). While the hydroxide cation $[9]^+$ persists long enough on the electrochemical time scale to be observed in the CV scans, the longer time scale of the chemical oxidation reaction with $FcPF_6$ permits a fluoride ion from the PF_6^- anion to displace the Zr-bound hydroxide ligand. Consistent with this hypothesis, the reaction of $[9]^+$ with $FcPF_6$ results in quantitative conversion to the fluoride species $[7]^+$. On the other hand, treating the $S = 1/2$ hydroxide compound **8** with stoichiometric $[^tBu_4N][PF_6]$ did not lead to any reaction, suggesting the important role of oxidation in OH^-/F^- exchange to generate the fluoride compound $[7]^+$. Additionally, a stoichiometric reaction between the hydroxide compound **8** and $FcPF_6$ affords a mixture of fluoride complex $[7]^+$ and hydroxide complex $[9]^+$, and a CV of this mixture highlights the distinguishable differences between their redox potentials (Fig. 7, green voltammogram).

The electronic structure of the oxidized species $[2]^+$ was investigated computationally using density functional theory (DFT) to better understand the instability of $[2]^+$ and the origin of its H-atom abstraction reactivity. DFT calculations were performed on two possible spin states of $[2]^+$, revealing the quartet to be $7.1 \text{ kcal mol}^{-1}$ more stable than the doublet spin state. In the quartet configuration, the unpaired electrons in $[2]^+$ were found to be predominately localized on the cobalt center, with a small but not negligible amount of spin density residing on the Zr-bound oxo ligand (Fig. 8A). The unpaired spin density on the oxo ligand results from a three-center four-electron bonding interaction involving the O, Zr, and Co atoms. As a result, the highest doubly occupied molecular orbital of **2** has both Co and O character (Fig. 8B), leading to unpaired electron density on the oxygen atom of $[2]^+$ upon oxidation of **2**. Since the Zr-bound oxygen atom is the least sterically protected site with significant spin density, the computational results provide an excellent explanation for the observed H-atom abstraction process that occurs upon oxidation of **2**.

Conclusions

In summary, the reactivity of the terminal oxo heterobimetallic Zr/Co compound **2** was explored, taking advantage of the steric protection of the mesityl ligand substituents that prevent oxo ligand bridging modes. In contrast to a monometallic Zr^{IV} oxo compound, **2** exhibited redox chemistry that was enabled by the pendent Co center, leading to the synthesis and isolation of a series of bimetallic Zr/Co compounds **3–9**. Although no direct metal–metal bond is present in **2**, outer-sphere redox changes mediated by the Co center are shown to work in concert with the inner-sphere nucleophilic reactivity of the reactive $Zr=O$ moiety. Both experimental and computational studies suggested that one-electron oxidation of **2** affords a transient oxidized species, $[2]^+$, containing a nucleophilic Zr oxo fragment and a tethered oxidizing cobalt center that work in concert to facilitate H-atom abstraction. This study represents another mechanism by which early/late heterobimetallic compounds can initiate unique reactivity *via* participation of two metals that play distinct roles. In this case, the late metal (Co) serves as



a redox-active metalloligand to mediate redox reactions at a d⁰ Zr^{IV}-oxo fragment.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This research was supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, Catalysis Science Program, under Award No. DE-SC0019179. The authors gratefully acknowledge Dr Alicia K. Friedman for assistance with characterization of several compounds using mass spectrometry. Caitlin M. Wade is acknowledged for assistance with preparation of the graphical abstract. The authors are also grateful for access to the Ohio Supercomputer Center.⁵⁰

Notes and references

- 1 K. B. Sharpless and T. C. Flood, *J. Am. Chem. Soc.*, 1971, **93**, 2316–2318.
- 2 J. S. M. Wai, I. Marko, J. S. Svendsen, M. G. Finn, E. N. Jacobsen and K. B. Sharpless, *J. Am. Chem. Soc.*, 1989, **111**, 1123–1125.
- 3 G. Cainelli and G. Cardillo, *Chromium Oxidations in Organic Chemistry*, Springer Berlin Heidelberg, Berlin, Heidelberg, 1984, vol. 19.
- 4 A. J. Fatiadi, *Synthesis*, 1987, 85–127.
- 5 P. R. Ortiz de Montellano, *Cytochrome P450: structure, mechanism, and biochemistry*, Springer International Publishing, 4th edn, 2015.
- 6 J. T. Groves, W. J. Kruper and R. C. Haushalter, *J. Am. Chem. Soc.*, 1980, **102**, 6375–6377.
- 7 M. Schroeder, *Chem. Rev.*, 1980, **80**, 187–213.
- 8 K. B. Wiberg and W. S. Trahanovsky, *Oxidation in organic chemistry*, Academic Press, 1965.
- 9 R. H. Holm and J. M. Berg, *Acc. Chem. Res.*, 1986, **19**, 363–370.
- 10 R. H. Holm, *Chem. Rev.*, 1987, **87**, 1401–1449.
- 11 A. Gunay and K. H. Theopold, *Chem. Rev.*, 2010, **110**, 1060–1081.
- 12 T. A. Betley, Q. Wu, T. Van Voorhis and D. G. Nocera, *Inorg. Chem.*, 2008, **47**, 1849–1861.
- 13 K. P. Kepp, *Inorg. Chem.*, 2016, **55**, 9461–9470.
- 14 G. Fachinetti, C. Floriani, A. Chiesi-Villa and C. Guastini, *J. Am. Chem. Soc.*, 1979, **101**, 1767–1775.
- 15 L. Rozes, N. Steunou, G. Fornasieri and C. Sanchez, *Monatshefte für Chemie*, 2006, **137**, 501–528.
- 16 P. Coppens, Y. Chen and E. Trzop, *Chem. Rev.*, 2014, **114**, 9645–9661.
- 17 F. A. Cotton, *Advanced inorganic chemistry*, 5th edn, 1988.
- 18 C. E. Housmeckerides, D. L. Ramage, C. M. Kretz, J. T. Shontz, R. S. Pilato, G. L. Geoffroy, A. L. Rheingold and B. S. Haggerty, *Inorg. Chem.*, 1992, **31**, 4453–4468.
- 19 P. A. Goodson and D. J. Hodgson, *Inorg. Chem.*, 1989, **28**, 3606–3608.
- 20 R. Guillard, J. M. Latour, C. Lecomte, J. C. Marchon, J. Protas and D. Ripoll, *Inorg. Chem.*, 1978, **17**, 1228–1237.
- 21 J. R. Hagadorn and J. Arnold, *Organometallics*, 1998, **17**, 1355–1368.
- 22 A. Bodner, P. Jeske, T. Weyhermueller, K. Wieghardt, E. Dubler, H. Schmalle and B. Nuber, *Inorg. Chem.*, 1992, **31**, 3737–3748.
- 23 A. J. Hoskin and D. W. Stephan, *Organometallics*, 1999, **18**, 2479–2483.
- 24 A. Mendiratta, J. S. Figueroa and C. C. Cummins, *Chem. Commun.*, 2005, 3403.
- 25 J. P. Krogman, M. W. Bezpalko, B. M. Foxman and C. M. Thomas, *Inorg. Chem.*, 2013, **52**, 3022–3031.
- 26 D. Jacoby, S. Isoz, C. Floriani, A. Chiesi-Villa and C. Rizzoli, *J. Am. Chem. Soc.*, 1995, **117**, 2793–2804.
- 27 M. R. Smith, P. T. Matsunaga and R. A. Andersen, *J. Am. Chem. Soc.*, 1993, **115**, 7049–7050.
- 28 H. Zhang, G. P. Hatzis, C. E. Moore, D. A. Dickie, M. W. Bezpalko, B. M. Foxman and C. M. Thomas, *J. Am. Chem. Soc.*, 2019, **141**, 9516–9520.
- 29 W. A. Howard, M. Waters and G. Parkin, *J. Am. Chem. Soc.*, 1993, **115**, 4917–4918.
- 30 M. J. Carney, P. J. Walsh and R. G. Bergman, *J. Am. Chem. Soc.*, 1990, **112**, 6426–6428.
- 31 M. J. Carney, P. J. Walsh, F. J. Hollander and R. G. Bergman, *J. Am. Chem. Soc.*, 1989, **111**, 8751–8753.
- 32 W. A. Howard and G. Parkin, *J. Am. Chem. Soc.*, 1994, **116**, 606–615.
- 33 I. G. Powers and C. Uyeda, *ACS Catal.*, 2017, **7**, 936–958.
- 34 P. Buchwalter, J. Rosé and P. Braunstein, *Chem. Rev.*, 2015, **115**, 28–126.
- 35 R. C. Cammarota, L. J. Clouston and C. C. Lu, *Coord. Chem. Rev.*, 2017, **334**, 100–111.
- 36 N. P. Mankad, *Chem.–Eur. J.*, 2016, **22**, 5822–5829.
- 37 J. P. Krogman, M. W. Bezpalko, B. M. Foxman and C. M. Thomas, *Dalton Trans.*, 2016, **45**, 11182–11190.
- 38 B. P. Greenwood, G. T. Rowe, C.-H. Chen, B. M. Foxman and C. M. Thomas, *J. Am. Chem. Soc.*, 2010, **132**, 44–45.
- 39 S. L. Marquard, M. W. Bezpalko, B. M. Foxman and C. M. Thomas, *Organometallics*, 2014, **33**, 2071–2079.
- 40 S. L. Marquard, M. W. Bezpalko, B. M. Foxman and C. M. Thomas, *J. Am. Chem. Soc.*, 2013, **135**, 6018–6021.
- 41 J. P. Krogman, B. M. Foxman and C. M. Thomas, *J. Am. Chem. Soc.*, 2011, **133**, 14582–14585.
- 42 C. M. Thomas, J. W. Napoline, G. T. Rowe and B. M. Foxman, *Chem. Commun.*, 2010, **46**, 5790.
- 43 H. Zhang, B. Wu, S. L. Marquard, E. D. Litle, D. A. Dickie, M. W. Bezpalko, B. M. Foxman and C. M. Thomas, *Organometallics*, 2017, **36**, 3498–3507.
- 44 J. W. Beattie, C. Wang, H. Zhang, J. P. Krogman, B. M. Foxman and C. M. Thomas, *Dalton Trans.*, 2020, **49**, 2407–2411.
- 45 J. W. Napoline, J. P. Krogman, R. Shi, S. Kuppaswamy, M. W. Bezpalko, B. M. Foxman and C. M. Thomas, *Eur. J. Inorg. Chem.*, 2013, **2013**, 3874–3882.
- 46 M. Gomberg, *J. Am. Chem. Soc.*, 1900, **22**, 752–757.



- 47 E. S. Jang, C. L. McMullin, M. Käß, K. Meyer, T. R. Cundari and T. H. Warren, *J. Am. Chem. Soc.*, 2014, **136**, 10930–10940.
- 48 N. G. Connelly and W. E. Geiger, *Chem. Rev.*, 1996, **96**, 877–910.
- 49 B. P. Greenwood, S. I. Forman, G. T. Rowe, C.-H. Chen, B. M. Foxman and C. M. Thomas, *Inorg. Chem.*, 2009, **48**, 6251–6260.
- 50 Ohio Supercomputer Center, 1987, Columbus, OH, <http://osc.edu/ark:/19495/f5s1ph73>.

