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Effect of σ^* P--O Orbital on Structure, Stereomutation, and Reactivity of C -Apical O -Equatorial Spirophosphoranes

Kin-ya Akiba^a, Shiro Matsukawa^b, Takahiro Adachi^b, Yohsuke Yamamoto^b, Suyong Re^c & Shigeru Nagase^c

^a Advanced Research Center for Science and Engineering, Waseda University, 3-4-1 Ohkubo, Tokyo, 169-8555, Japan

^b Department of Chemistry, Graduate School of Science, Hiroshima University, 1-3-1 Kagamiyama, Higashi-Hiroshima, 739-8526, Japan

^c Department of Theoretical Studies, Institute for Molecular Science, Myodaiji, Okazaki, 444-8585, Japan

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EFFECT OF $\sigma^*_{P=0}$ ORBITAL ON STRUCTURE, STEREOMUTATION, AND REACTIVITY OF C-APICAL O-EQUATORIAL SPIROPHOSPHORANES

Kin-ya Akiba,^a Shiro Matsukawa,^b Takahiro Adachi,^b Yohsuke Yamamoto,^b Suyong Re,^c and Shigeru Nagase^c Advanced Research Center for Science and Engineering, Waseda University, 3-4-1 Ohkubo, Tokyo 169-8555, Japan;^a Department of Chemistry, Graduate School of Science, Hiroshima University, 1-3-1 Kagamiyama, Higashi-Hiroshima 739-8526, Japan;^b and Department of Theoretical Studies, Institute for Molecular Science, Myodaiji, Okazaki 444-8585, Japan^c

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Effect of σ^*_{P-O} orbital of C-apical O-equatorial (O-cis) spirophosphorane was investigated both experimentally and theoretically. O-cis phosphoranes revealed to be much more electrophilic on the phosphorus atom than O-trans isomers by experimental studies. Theoretically, the energy of the σ^*_{P-O} orbital of O-cis phosphorane was calculated to be lower than that of the σ^*_{P-C} orbital of O-trans phosphorane by 18.7 kcal/mol, and the result supports the enhanced electrophilicity of O-cis spirophosphoranes compared with O-trans isomer.

Keywords: Hypervalent; $\sigma^*{}_{\rm P\!-\!O}$ orbital; phosphorus; reactivity; reversed apicophilicity

INTRODUCTION

We have reported on the synthesis, structure, and stereomutation of *C*-apical *O*-equatorial (*O*-*cis*) spirophosphorane **1a** and its stable stereoisomer of *O*-apical *C*-equatorial (*O*-*trans*) spirophosphorane **2a**.¹ The phosphorane **1a** was the first example exhibiting reversed apicophilicity, which could be converted to its stable pseudorotamer **2a**. *O*-*cis* phosphorane **1** has one of the two P–O bonds on the equatorial plane and therefore has a σ^*_{P-O} orbital on the same plane. On the other

Address correspondence to K.-y. Akiba, Advanced Research Center for Science and Engineering, Waseda University, 3-4-1 Ohkubo, Tokyo 169-8555, Japan. E-mail: akibaky@waseda.jp



FIGURE 1

hand, *O*-trans phosphorane **2** has the corresponding P–C bond on the equatorial plane; hence σ^*_{P-C} , too (Figure 1). Since the energy level of σ^*_{P-O} should be significantly lower than that of σ^*_{P-C} , there should be an apparent effect of σ^*_{P-O} to show the difference between **1** and **2** in reactivity such as nucleophilicity and stability of α -carbanions. Now we describe the effects of σ^*_{P-O} orbital of **1** from both an experimental and theoretical viewpoint.

RESULTS AND DISCUSSION

Synthesis of *O-cis* Spirophosphoranes by Oxidative Cyclization of the Dianion

As reported in 1996, we succeeded to isolate *O*-cis spirophosphorane **1a** by thermal cyclization of P–H(apical) monocyclic phosphorane.¹ However this method is not suitable for synthesis of *O*-cis phosphoranes because the thermal conditions accerelate stereomutation to the corresponding *O*-trans isomer. Indeed, **1a** was obtained in 71% yield and **2a** in 29% yield by this thermal method. To improve this problem, a milder oxidative cyclization method was developed in our laboratory² (Scheme 1). The new protocol provided **1a** and **1b** exclusively at low temperatures in high yields, and aryl derivative **1c** could also be synthesized for the first time.

The possible reaction mechanism is shown in Scheme 2. The reaction of P–H (equatorial) phosphorane **A** with more than 2 equivalents of RLi provides the dianion **B**, where one equivalent of RLi works as a base and the other as a nucleophile. In the resulting dianion **B**, the lone pair electrons on phosphorus occupy an equatorial position because of its extreme electron-donicity. In this situation, oxidant I_2 reacts with the



dianion **B** to produce P–I (equatorial) phosphorane **C**, and the remaining alkoxide anion rapidly cyclizes with simultanious elimination of I^- to form *O*-*cis* spirophosphorane **1** exclusively.

Enhanced Electrophilicity of *O-cis* Phosphorane 1: Effect of $\sigma^*_{P=0}$ Orbital in the Equatorial Plane

Reactions of *O-cis* **1a** and *O-trans* **2a** with nucleophiles were examined. Using TBAF (tetrabutylammonium fluoride; $[n-\text{Bu}]_4\text{N}^+\text{F}^-$) as a nucleophile, the reaction of **1a** readily afforded a hexacoordinate phosphate bearing a P–F bond **3** (${}^1J_{\text{P}-\text{F}} = 706 \text{ Hz}$) while **2a** did not react at all. Configuration of the phosphate **3** could not be determined because of rapid decomposition by trace amounts of H₂O. However, we have already characterized the corresponding hexacoordinate fluoroantimonate with two Martin ligands by X-ray analysis;³ therefore, the fluorine atom in phosphate **3** is also likely to be located anti to the oxygen due to the trans influence as the fluoroantimonate. In the case of the reaction with MeLi, **1a** reacted under mild conditions to afford a monocyclic phosphorane **4** after aqueous work up but **2a** did not (Scheme 3).



Stabilization of Lone Pair Electrons Adjacent to the Phosphorus of *O-cis* 1: Effect of $\sigma^*_{\rm P-O}$ Orbital in the Equatorial Plane

Deprotonation of *O-cis* **1d** with KHMDS (potassium hexamethyldisilazide; $(Me_3Si)_2N^-K^+$) occurred in THF at 0°C during 30 min and gave α -deuterated product (40% yield) after quenching the mixture with D₂O. However, *O-trans* **2d** could not be deprotonated under the same conditions. Moreover, the benzyl anion of **1d** was much more stable thermally than the neutral molecule of **1d**. In THF, *O-cis* **1d** was totally converted to *O-trans* **2d** at 60°C during 5 h. Under the same conditions, the benzyl anion of **1d** remained almost unchanged, which shows the significant stability of the *O-cis* benzyl anion compared to the corresponding neutral molecule.

Theoretical Study on σ^* Orbitals of *O-cis* and *O-trans* Phosphoranes in the Equatorial Plane

In order to evaluate a difference in energy between the σ^*_{P-O} orbital of *O*-*cis* phosphorane and that of *O*-*trans* isomer, theoretical calculations for **1b** and **2b** were carried out (B3LYP/6-31G(d)). As shown in Figure 2, the energy of σ^*_{P-O} of **1b** (LUMO+4) is lower than that of σ^*_{P-C} of **2b** (LUMO+5) by 18.7 kcal/mol. Other LUMOs below σ^*_{P-X} (X = O or C) were distributed on the aromatic rings of the Martin ligands. Therefore, the responsible orbitals toward nucleophilicity and stabilization of α -carbanion should be the σ^*_{P-X} orbitals. As a result, the large



FIGURE 2

difference in energy between σ^*_{P-O} and σ^*_{P-C} orbital provided theoretical evidence for the enhanced reactivity of *O*-*cis* spirophosphorane **1** compared with *O*-*trans* isomer **2**, as shown in Scheme 3.

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