Photooxidation of Mixed Aryl and Biarylphosphines

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ABSTRACT

Arylphosphines and dialkylbiarylphosphines react with singlet oxygen to form phosphine oxides and phosphinate esters. For mixed arylphosphines, the most electron-rich aryl group migrates to form the phosphinate, while for dialkylbiarylphosphines migration of the alkyl group occurs. Dialkylbiarylphosphines also yield arene epoxides, especially in electron-rich systems. Phosphinate ester formation is increased at high temperature, while protic solvents increase the yield of epoxide. The product distribution provides evidence for Buchwald's recent conformational model for the aerobic oxidation of dialkylbiarylphosphines.

Phosphadioxiranes are highly reactive heteroatom-containing peroxides and belong to what Greer et al. have aptly called "rather exotic types of cyclic peroxides".¹ It is now wellestablished that they are the primary intermediates in the reaction of singlet dioxygen with phosphines.^{2,3} In addition, at very high temperatures, phosphadioxiranes are probably formed during the reaction of triplet dioxygen with phosphines.⁴ Intermolecular oxygen atom transfer from these highly unstable intermediates to the starting phosphine leads to phosphine oxide formation. In addition to this process, phosphadioxiranes may undergo a variety of interesting

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reactions, including oxygen atom insertion into the aryl P-^C bond to form phosphinate esters and, in a few cases, arene epoxidation.^{5,2b,c} Several theoretical studies suggest that phosphadioxiranes should be electrophilic oxidants.³ However, the scope of oxygen atom transfer reactions from phosphadioxirane as well as the regioselectivity and effects of temperature and solvent are not known. For example, the photooxidation of arylphosphines with different aryl groups attached to the phosphorus atom has not been studied; i.e., it is not known if the formation of phosphinate esters is regioselective. Furthermore, there have been no reports of photooxidation of several other important classes of phosphines such as phosphines bearing biaryl ligands. Intramolecular oxygen atom insertion into the P-C bond and arene epoxidation might ultimately be useful reactions for the functionalization of such phosphines, if the scope and regioselectivity of these reactions were known, and if the competing intermolecular oxygen atom transfer to form phosphine oxide could be limited. In this paper, we present a detailed investigation of product distribution, regioselectivity, solvent, and temperature effects in the reactions of phosphadioxiranes generated from singlet oxygen and a variety of different phosphines. We show that oxygen atom insertion in mixed (i.e., bearing different aryl and/or alkyl groups) phosphines is, in fact, regioselective. We also investigate how solvent and temperature effects can be used to maximize the yield of either phosphinate ester or arene epoxide.

Photooxidation of arylphosphines bearing large substituents in the ortho position of the aryl ligand predominately yield phosphinate esters from intramolecular oxygen atom insertion into the $P-C$ bond.^{2b} Arylphosphines without bulky substituents give mainly phosphine oxide upon reaction with singlet oxygen. The intermediate phosphadioxirane is presumed to be an electrophilic oxidant, but there is little experimental evidence, as phosphines bearing different aryl ligands have not been studied. We have therefore begun our study of the photooxidation of mixed phosphines by investigating the reaction of bis(*o*-methoxyphenyl)phenylphosphine (**1**) with singlet oxygen (Scheme 1). Photooxidation

of 1 at room temperature (solvent $=$ toluene- d_8 , sensitizer $=$ tetraphenylporphyrin (TPP), phosphine concentration $=$ ¹-20 mM, tungsten-halogen lamp, cutoff filter at 493 nm) leads to only two products, namely (*o*-methoxyphenyl)phenyl *o*-methoxyphenylphosphinate (**2**), and bis(*o*-methoxyphenyl)phenylphosphine oxide (**3**).

Both products were characterized by ${}^{1}H$ and ${}^{31}P$ NMR, and their identities were confirmed by X-ray molecular structures (see the Supporting Information). The insertion reaction leading to formation of **2** is completely regioselective. We did not observe any oxygen-atom insertion into the ^P-C bond bearing the unsubstituted phenyl ring. Furthermore, unlike for binaphthylphosphines,^{2c} we did not observe any hydroxylation of the aromatic ring.

Dialkylbiarylphosphines have been widely used in Pdcatalyzed cross-coupling reactions, in part because they are rather resistant toward oxidation.^{6,7} Buchwald et al. have recently suggested that the biaryl ligand as well as substituents on the 2′ and 6′ positions of the biaryl ring play a key role in the susceptibility-or lack thereof-toward reaction with triplet oxygen.⁶ The reactivity of these phosphines with singlet oxygen has not been studied to date. A priori, a wide range of possible products can be envisioned, i.e., oxygen atom insertion into the P-C bond of either the alkyl or biaryl ligand, formation of arene epoxides, and simple formation of phosphine oxide by intermolecular oxygen atom transfer. Of particular interest is the question of whether alkyl or aryl group migration is preferred in these compounds; alkyl migration in a phosphadioxirane has not previously been reported.

The reaction of unsubstituted 2-di-*tert*-butylbiphenylphosphine (**4**) with singlet dioxygen in deuterated toluene at room temperature yields the corresponding phosphine oxide (**5**) as the sole product (see Table 1). No epoxidation or insertion

 α k_T values are averages of three runs; error is one standard deviation. ^{*b*} All reactions were performed in an NMR tube using 1 mL solutions of biphenylphosphine (24 mM) in toluene- d_8 , concentration of TPP sensitizer $= 0.13$ mM. Relative amounts of products were calculated (average of three runs, error is one standard deviation) using 31P NMR integrations.

products (i.e., via alkyl or aryl migration) were observed. Even at high temperature where aryl migration may be more favorable (see below), the reaction of singlet oxygen with biphenylphosphine **4** in toluene yields phosphine oxide **5** as the only product.

In marked contrast to the unsubstituted 2-di-*tert*-butylbiphenylphosphine **4**, the bulky highly substituted di-*tert*butyltriisopropylbiphenylphosphine **6** reacts with singlet

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dioxygen to form phosphine oxide **7** and phosphinate **8** (see Scheme 2), as well as a very small amount of arene epoxide **9** in toluene- d_8 . The latter compound is readily identified by the ¹ H NMR signals of the vinyl protons of the oxidized ring which give rise to singlets at 5.70 and 4.70 ppm. Even at room temperature, at 24 mM starting concentration of **6**, the phosphinate product **8** predominates (Table 1). The phosphinate ester **8** is of special interest: We were able to establish that migration in phosphines containing both alkyl and aryl ligands, migration of the alkyl group is favored over the aryl group. The ¹H NMR spectrum of a purified sample of **8** shows that the hydrogens of the *tert*-butyl group directly attached to the phosphorus are split by the phosphorus atom into a doublet $(J = 15.9 \text{ Hz})$ at 0.97 ppm. The hydrogen atoms of the *tert*-butyl group bound to the oxygen, on the other hand, exhibit no coupling to the phosphorus atom and thus occur as a singlet at 1.42 ppm. In contrast to phosphinate **8**, the hydrogens of both *tert*-butyl groups of phosphine oxide **7** are chemically equivalent and appear as a doublet $(J =$ 13.3 Hz) at 1.25 ppm. Additional details of the spectroscopic analyses are given in the Supporting Information.

Interestingly, the reactivity of the related dicyclohexyltriisopropylbiphenylphosphine **10** is rather different. During photooxidation of **10** at room temperature, formation of phosphine oxide **11** predominates relative to phosphinate ester **12**; i.e., intermolecular oxygen atom transfer is much more facile than for the di-*tert*-butylphosphine **6**. Furthermore, we observed a considerable amount of epoxide **13** (22% at room temperature for initial concentration of **10** of 24 mM, Table 1). The epoxide could be formed either by intramolecular or intermolecular oxygen atom transfer from the intermediate phosphadioxirane. If the epoxidation proceeds by an intramolecular pathway, the product ratio of phosphine oxide and epoxide vs starting phosphine concentration (at low conversion of phosphine) can be expressed by eq 1, where *k*epox is the rate constant for epoxide formation from phosphadioxirane and *k*^o is the rate constant for intermolecular oxygen atom transfer to give 2 equiv of phosphine oxide:

$$
\frac{[11]}{[13]} = \frac{2k_0[10]}{k_{\text{epox}}} \tag{1}
$$

A plot of the product ratio **11**/**13** vs **10** gave a straight line with a slope $(2k_o/k_{\text{epox}})$ of 110 M^{-1} , confirming that epoxidation is indeed an intramolecular process. The relative rate ratio of intermolecular oxygen atom transfer vs intramolecular epoxidation is 55 M^{-1} , and intermolecular oxygen atom transfer predominates, except at very low concentrations of **10** (2 mM or less).

Even though the photooxidation product distribution for phosphines **6** and **10** is very different, we determined that their rate constants for singlet oxygen removal k_T are very similar (Table 1, second column). This implies that the approach of the singlet oxygen molecule to phosphines **6** and **10** is not affected by the difference of the alkyl ligands. Hence the origin of the different product distribution results from steric factors of the intermediate phosphadioxirane complexes and their interaction with starting phosphines. Buchwald and Barder recently noted that phosphine **6** is considerably more resistant to reaction with ground state (triplet) oxygen than phosphine **10**. ⁶ They investigated the barrier for rotation around the $P-C$ bond of the biaryl ligand such that the phosphorus lone pair (or peroxidic intermediate) points away from the large biphenyl ligand. For compound **6**, this rotational barrier is much larger than for compound 10 (12.6 vs 6.3 kcal/mol).⁶ Furthermore, they determined that the "away conformer" (where the phosphorus lone pair or the dioxygen moiety of a peroxidic intermediate points away from the bulky biaryl ring system) is much more disfavored for compound **6** relative to compound **10**. Our observations for the product distribution during the photooxidation of **6** and **10** provide strong evidence for Buchwald's model. Since compound **6** has a much more restricted rotation and the "away conformer" is more strongly disfavored, intermolecular oxygen atom transfer to unreacted starting material is much more difficult, and intramolecular oxygen atom insertion predominates.

Replacement of the isopropyl groups of the biarylphosphines with isopropoxy groups also has a drastic effect on the reactivity of the intermediate phosphadioxirane: reaction of diisopropoxybiphenylphosphine 14 with singlet oxygen in toluene- d_8 gives epoxide **15** in nearly quantitative yield (Scheme 3).

Epoxide **15** is stable at 0 °C for weeks but decomposes into a complex mixture at room temperature. The rate of singlet oxygen removal (k_T) by this phosphine is faster

compared to the other biaryl phosphines, due to the electrondonating character of the isopropoxy groups.

The product distribution in all of the photooxidation reactions reported here is highly sensitive to temperature and solvent effects. In general, at higher temperature, the yield of phosphinate ester is increased and that of phosphine oxide is diminished. Details are given in Table 2. Thus, if the phosphinate esters

Table 2. Product Distribution during Photooxidation of Dialkylbiarylphosphines at Different Temperatures*^a*

of biphenylphosphine (24 mM) in toluene-*d*8. Relative amounts of products were calculated (average of three runs, error is one standard deviation) using ³¹P NMR integrations.

are desired products, carrying out the photooxidation at high temperature and low starting material concentrations will result in the highest yields of phosphinate esters.

The presence of protic solvents leads to protonation of the intermediate phosphadioxirane and formation of hydroperoxyphosphoranes.^{2a,b,3b} Very little is known about the chemistry of the latter species. To investigate the reactivity of the hydroperoxyphosphorane, we added up to 20% (by volume) of CD3OD to phosphines **6** and **10** prior to carrying out the photooxidation. Under these conditions, phosphinate ester formation is suppressed while the yield of epoxide and phosphine oxide is increased (Table 3). Clearly,

of biphenylphosphine (24 mM). Relative amounts of products were calculated (average of three runs, error is one standard deviation) using 31P NMR integrations.

the hydroperoxyphosphorane is much more capable than phosphadioxirane to form arene epoxides. Hence, if the latter product is desired, one can simply carry out the photooxidation under protic conditions. In general, despite the fact that photooxidation of biarylphosphines may lead to multiple products, judicious choice of temperature and solvent system can maximize the yield of one of the oxidation products over the others.

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Supporting Information Available: Experimental procedures, full spectroscopic data for all new compounds, and X-ray data for compounds **2** and **3** (CIF). This material is available free of charge via the Internet at http://pubs.acs.org.

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