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Radical-initiated P,P-metathesis reactions of diphosphanes: evidence from experimental and computational studies†

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By combining the diphosphanes Ar_2P-PAr_2 , where $Ar = C_6H_5$, $4-C_6H_4Me$, $4-C_6H_4OMe$, $3,5-C_6H_3(CF_3)_2$, it has been shown that P,P-metathesis generally occurs rapidly under ambient conditions. DFT calculations have shown that the stability of unsymmetrical diphosphanes Z_2P-PZ_2' is a function of the difference between the Z and Z' substituents in terms of size and electronegativity. Of the mechanisms that were calculated for the P,P-metathesis, the most likely was considered to be one involving Ar_2P^* radicals. The observations that photolysis increases the rate of the P,P-metatheses and TEMPO inhibits it, are consistent with a radical chain process. The P,P-metathesis reactions that involve $(o-Tol)_2P-P(o-Tol)_2$ are anomalously slow and, in the absence of photolysis, were only observed to take place in CHCl₃ and CH₂Cl₂. The role of the chlorinated solvent is ascribed to the formation of Ar_2PCl which catalyses the P,P-metathesis. The slow kinetics observed with $(o-Tol)_2P-P(o-Tol)_2$ is tentatively attributed to the $o-CH_3$ groups quenching the $(o-Tol)_2P^*$ radicals or inhibiting the metathesis reaction sterically.

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Introduction

Dynamic and reversible chemical reactions have garnered increasing attention in medicine and biology, supramolecular chemistry and materials science, and they are key to the rapidly growing field of dynamic covalent chemistry (DCC). 1,12,13 In DCC, disulfides play a leading role, by virtue of the weakness of the S–S bond and their ready availability *via* chemical synthesis and their presence in natural products. 12,14-16 In the last decade, several other main group element–element bonds (O–O, 17,18 N–N 19 and Se–Se 9,20,21) have been investigated for their potential in dynamic chemistry with a view to their application in the field of polymeric materials. A notable omission from this group is diphosphanes (P₂R₄) which contain a P–P bond whose strength (*ca.* 52 kcal mol⁻¹)²² is intermediate between S–S (*ca.* 62 kcal mol⁻¹)²³ and Se–Se (*ca.* 40 kcal mol⁻¹)²² and therefore diphosphanes should be candidates for DCC.

Diphosphanes have received considerable attention as targets themselves and also as undesired byproducts in the syn-

thesis of tertiary phosphines. 24-31 Several methods to prepare diphosphanes are available, including: Wurtz-type reductive coupling of chlorophosphines (with Li, Na, K, Mg and Hg);³²⁻³⁴ salt metathesis between chlorophosphine and LiPR2; 26,27 dehydrocoupling of HPR₂;³⁵⁻³⁹ chlorosilane elimination;⁴⁰ P-N/P-P bond metathesis.41 The chemistry of diphosphanes can be divided into two categories: reactions where the P-P bond remains intact and reactions where the P-P bonds are cleaved. 42 Reactions involving P-P cleavage are pertinent to the chemistry described in this article and include additions of X2 to the R2P- PR_2 to give monophos species R_2PX (e.g. X = H, Cl) and additions of R_2P-PR_2 to C=C or C=C bonds (diphosphination) to give diphos species (Scheme 1). It has also been shown that diphosphanes containing P-N bonds can add to C=S or C=O bonds. 43,44 Mechanisms have been proposed for some of the P-P cleavage reactions including radical chain processes. 43,45

Diphosphane metathesis has not been systematically investigated, but an early ³¹P NMR study by Harris *et al.* showed that rapid metathesis occurred between Me₂P-PMe₂ and (F₃C)₂P-P(CF₃)₂ in CH₂Cl₂ to form Me₂P-P(CF₃)₂. ^{46,47} Gilheany *et al.* ⁴⁸ have shown that the calculated high energy barrier to P,P-metathesis *via* a concerted mechanism is not compatible with the observed rapid kinetics for the Me₂P-P(CF₃)₂ system and suggested that pathways involving impurities in the diphosphanes may facilitate the P,P-metathesis process. Grubba *et al.* ²⁶ have succeeded in the preparation of a variety of pure, unsymmetrical diphosphanes containing P-C(alkyl) and P-N bonds. By

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Scheme 1 Some reactions involving P-P cleavage of tetraphenyldiphosphane. 42

$$X_2$$

Scheme 2

employing low temperatures, they avoided unwanted P.P-metatheses that would lead to symmetrical diphosphanes contaminating the products. Moreover, they reported that R₂PCl or R₂PLi can catalyse diphosphane metathesis. We have reported that diphosphane metathesis was a side reaction in the attempted synthesis of some unsymmetrical diphosphanes.²⁷

In this study, we have investigated the P,P-metathesis reactions of tetra-aryldiphosphanes (Scheme 2) using ³¹P{¹H} NMR spectroscopy and computational methods, with a view to exploring the dynamic nature of these P-P bonds. This has provided some new insights into the diphosphane metathesis reaction.

Results and discussion

Thermodynamics of diphosphane metathesis

Experimental equilibrium studies. The tetra-aryldiphosphanes A2-E2 (Chart 1) used in this study, were chosen in order to sample a range of steric and electronic effects. All have been previously reported and were prepared by modified literature procedures for the reductive coupling of the corresponding chlorophosphines by Mg in THF (see ESI† for details).26,34 In each of the diphosphane samples, residual chlorophosphine was not detected by 31P NMR spectroscopy and was therefore estimated to be less than 0.05% (see ESI†).

When two of the diphosphanes A₂-E₂ were mixed in CDCl₃ at ambient temperature, a reaction ensued to give an equilibrium mixture of homodiphosphanes X2 and Y2, and the heterodiphosphane XY (Scheme 2). The data for the diphosphane metathesis reactions are given in Table 1.

The ³¹P{¹H} NMR data for the XY species (Table 1) show that the chemical shifts for the two signals are close to the respective X_2 and Y_2 with ${}^1J_{XY} \simeq 160$ Hz. For example, Fig. 1 is

the ³¹P{¹H} NMR spectrum obtained after mixing diphosphanes B2 and C2 (entry 4, Table 1) and shows the singlets for the reactants along with an AB pattern for the product BC; the spectra for the other diphosphane combinations are given in the ESI.†

The pure heterodiphosphane AE was prepared in order to establish the P,P-metathesis equilibrium with A2 and E2 (Scheme 2) from the opposite direction. Grubba et al. 26 have prepared a range of heterodiphosphanes (Z₂P-PZ'₂) by the addition of Z₂PLi to Z'₂PCl at low temperatures. However, our attempts to extend this route to the preparation of AE were unsuccessful, even with the reaction mixture maintained at -78 °C throughout; instead, an equilibrium mixture of A₂, E₂ and AE was obtained. This contrasting behaviour may be due to the lack of thermodynamic stability of heterodiphosphane **AE** compared with the Grubba heterodiphosphanes²⁶ (see below). The desired AE was prepared in >96% purity (after recrystallisation from methanol) via the protection/deprotection method (Scheme 3) that we previously developed for the synthesis of heterodiphosphanes. 27,49 Solutions of pure AE in CDCl3 do indeed equilibrate to a mixture of A2, E2 and AE (see below for further details).

The diphosphane equilibrium constants for the P,P-metatheses (Scheme 2) fall in the range 3-30 (see Table 1), which corresponds to ΔG values between -0.5 and -2 kcal mol⁻¹, demonstrating that the diphosphanes in this study are very close in energy and insensitive to the steric and electronic effects of the aryl substituents. The equilibria that do not involve E2 were established rapidly (see Table 1) despite the dilution (12.5 mM) which is a useful property for the potential application of diphosphanes in dynamic chemistry. The anomalously slow P,P-metatheses involving E2 are discussed later.

Chart 1

 Table 1
 Data for the P,P-metathesis equilibria shown in Scheme 2

Entry	X_2	\mathbf{Y}_2	K	Time ^a /min	³¹ P NM	R data fo	or XY
1^b	\mathbf{A}_2	\mathbf{B}_2	4	30	-14.9	-16.4	$J_{\rm PP}$ = 162 Hz
2^c	$\mathbf{A_2}$	C_2	5	20	-15.9	-18.4	$J_{\rm PP}$ = 159 Hz
3^d	$\mathbf{A_2}$	$\mathbf{D_2}$	30^e	90	-15.9	-8.5	$J_{\rm PP}$ = 177 Hz
4^d	$\mathbf{B_2}$	C_2	4	15	-16.8	-18.8	$J_{\rm PP}$ = 156 Hz
5^b	$\mathbf{B_2}$	$\mathbf{D_2}$	15	20	-11.7	-15.7	$J_{\rm PP}$ = 165 Hz
6^b	C_2	$\mathbf{D_2}$	3	20	-10.5	-15.4	$J_{\rm PP}$ = 173 Hz
7^c	$\mathbf{A_2}$	$\mathbf{E_2}$	4	700	-16.6	-35.3	$J_{\rm PP}$ = 152 Hz
8 ^c	$\mathbf{B_2}$	$\mathbf{E_2}$	4	1000	-19.1	-36.2	$J_{\rm PP}$ = 149 Hz
9^c	C_2	$\mathbf{E_2}$	4	2800	-19.0	-36.9	$J_{\rm PP}$ = 148 Hz
10^c	\mathbf{D}_2	$\mathbf{E_2}$	30	4000	-17.9	-29.5	J_{PP} = 156 Hz

 a Time by which equilibrium had been established according to 31 P NMR spectroscopy. b In THF. c In CDCl $_3$. d In C $_6$ D $_6$. e Estimated value (see details in ESI †).

Computational equilibrium studies. To understand better the factors that control the stability of heterodiphosphanes with respect to the corresponding homodiphosphanes, DFT calculations were carried out to determine the thermodynamics of the hypothetical equilibria shown in Scheme 4 in which the range of P-substituents Z and Z' is greatly extended from the substituted aryls in Scheme 2. The values for ΔG are collected in Table 2, and for some of the combinations (Table 2, entries 1–3 and 7–10), the calculated $|\Delta G|$ is less than 3 kcal mol⁻¹ and therefore, these particular P,P-metathesis equilibria are predicted to be finely balanced. The calculated

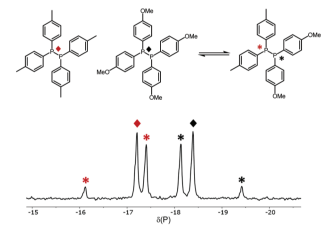


Fig. 1 The $^{31}P\{^{1}H\}$ NMR spectrum of the P,P-metathesis reaction between $(p-\text{Tol})_{2}P-P(p-\text{Tol})_{2}$ (B₂) and $(p-\text{Anisyl})_{2}P-P(p-\text{Anisyl})_{2}$ (C₂).

Scheme 3

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Table 2 Calculated ΔG values for the diphosphane metathesis equilibria Scheme 4

Entry	Z	\mathbf{Z}'	$\Delta G/\text{kcal mol}^-$
1	Me	Н	-0.5
2	Me	NH_2	-2.9
3	Me	OH	-2.9
4	Me	OMe	-3.7
5	Me	F	-6.6
6	Me	CF_3	-10.5
7	Ph	H	0.8
8	Ph	Me	-0.7
9	Ph	NH_2	1.3
10	Ph	NMe_2	-0.8
11	Ph	OMe	-4.7
12	Ph	F	-6.1
13	Ph	CF_3	-4.4
14	t Bu	Н	-13.4
15	^t Bu	Me	-3.3
16	^t Bu	Ph	-4.0
17	t Bu	NH_2	-6.8
18	^t Bu	NMe_2	-6.8
19	^t Bu	OMe	-11.2
20	^t Bu	\mathbf{F}	-15.4
21	^t Bu	CF_3	-3.4
22	o-Tol	Ph	1.2

 ΔG of 1.2 kcal mol⁻¹ (Table 2, entry 22) for $\mathbf{A}_2/\mathbf{E}_2$ is remarkably close to the experimental value of -0.8 kcal mol⁻¹ obtained from the experimentally determined value of K=4 in CDCl₃ solution (Table 1, entry 7).

Two gross trends emerge from the data in Table 2: (1) a bias in favour of the XY species is calculated when there is a large electronegativity difference between the X and Y fragments; thus Me₂P-P(CF₃)₂ is predicted (and experimentally observed⁵⁰) to be highly favoured (Table 2, entry 6); (2) a strong bias in favour of the XY species is calculated when there is a large difference in steric bulk between the X and Y fragments; thus ^tBu₂P-PH₂ is predicted to be highly favoured (Table 2, entry 13). The largest calculated ΔG in favour of the XY species is for ^tBu₂P-PF₂ (Table 2, entry 20), at an apparent apotheosis of both trends. The observed trends in the calculated relative stabilities of the XY species in Scheme 4 can be rationalised as follows: (a) electronegative Z substituents (such as F) will lead to repulsive electrostatic interactions between the δ + charges generated on the adjacent P atoms in diphosphane X2; (b) bulky Z' substituents (such as ^tBu) will lead to steric congestion in diphosphane Y2; (c) in XY, both of the P-P bond-destabilising effects identified in (a) and (b) will be reduced and, in some cases, a destabilising, repulsive, electrostatic $(\delta + / \delta +)$ interaction is replaced by a stabilising, attractive, electrostatic $(\delta + / \delta -)$ interaction.

The calculated ΔG values for some of the Grubba heterodiphosphanes²⁶ (Table 2, entries 16 and 18) suggest that these should have high thermodynamic stability with respect to the corresponding homodiphosphanes. By contrast, the heterodiphosphine **AE**, as reported above, is calculated to be of similar energy to the homodiphosphanes **A**₂ and **E**₂ (Table 2, entry 22).

Computational mechanistic studies of diphosphane metathesis

The mechanism of the diphosphane metathesis shown in Scheme 5 has been previously investigated computationally by Gilheany *et al.*⁴⁸ The lowest energy route that they could find involved the ionic intermediate shown in Scheme 5 but the calculated barrier of 36.5 kcal mol⁻¹ was too high to explain the rapidity of the P,P-metathesis that is observed experimentally.

We have investigated the mechanism of diphosphane metathesis computationally using the degenerate exchange reactions of Me_2P-PMe_2 (Scheme 6). Tetramethyldiphosphane was chosen in order to limit the number of rotamers and thus the conformational noise in the calculations. It was hoped to distinguish between the three plausible pathways (i–iii) shown in Scheme 7: (i) a [2+2] concerted process involving a 4-mem-

Scheme 5 Gilheany mechanism for a P,P-heterometathesis.

(i)
$$Me_2P - PMe_2$$
 $Me_2P - PMe_2$ $Me_2P - PMe_2$ $Me_2P - PMe_2$ $Me_2P - PMe_2$ (ii) $Me_2P - PMe_2$ $Me_$

Scheme 7

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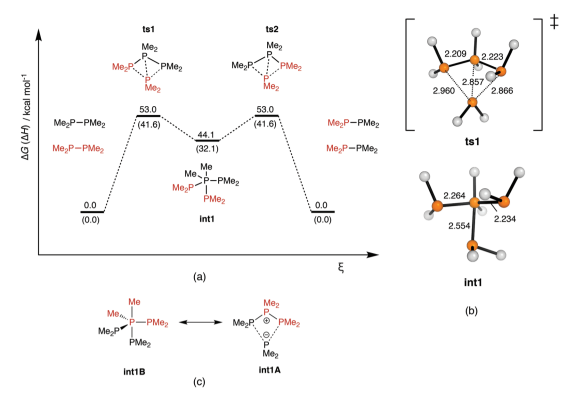


Fig. 2 (a) Reaction free energy profile for the P,P-homometathesis of Me₂P-PMe₂. Calculations performed at the PBE0-D3BJ/def2-TZVPP/PBE0-D3BJ/def2-SVP level of theory. (b) Calculated structures of ts1 and int1. Note: ts1 and ts2 are the same; key distances given in Å. (c) Resonance forms of int1

bered intermediate or transition state; (ii) nucleophilic attack by one diphosphane on another to give an ionic intermediate (similar to the one suggested in Scheme 5); (iii) a [2 + 1] radical chain process.

The overall reaction profile calculated for the thermal rearrangement of Me₂P-PMe₂ (shown in Fig. 2(a) proceeds via intermediate int1 depicted in its lowest energy conformation in Fig. 2(b). The structure of int1 can be viewed in terms of the two resonance forms int1A and int1B shown in Fig. 2(c) (further details are given in the ESI†). The ionic form int1A is analogous to the intermediate suggested for the P,P-heterometathesis shown in Scheme 5 and the covalent int1B contains an unusual central P(V) moiety. The profile shown in Fig. 2 corresponds to pathway (ii) of Scheme 7, and notably, attempts to locate a synchronous, σ-bond metathesis pathway (corresponding to pathway (i) in Scheme 7) were unsuccessful; no imaginary normal modes corresponding to the 2 bond-breaking/ forming vibrations are present along the relaxed, symmetric P4 ring expansion surface. The energy of **int1** of 44.1 kcal mol⁻¹ for the PMe2 species is lower for the analogous PPh2 intermediate $(\Delta G = 38.1 \text{ kcal mol}^{-1}, \text{ see ESI}^{\dagger})$ but this is calculated to be not lowered by solvation with CH2Cl2. Thus, the calculated very high barrier is incompatible with the rapidity of the Ar₂P-PAr₂ metathesis reactions that is observed experimentally.

The radical chain pathway (iii) in Scheme 7 requires homolysis of a P-P bond. Thermolysis is not viable under ambient conditions because the calculated P-P bond enthalpy is of the

order of 52 kcal mol⁻¹. However, DFT calculations for Ph_2P-PPh_2 (A_2) suggest that excitation from $S_0 \rightarrow S_1$ or $S_0 \rightarrow S_2$ would result in rapid homolysis of the P-P bond, yielding phosphanyl radicals (Ar_2P^*) *via* intersystem crossing from S_1 or S_2 to the first excited triplet state (T_1) of A_2 ; the T_1 state is dissociative, and would lead to spontaneous radical formation. From the Ar_2P^* , the P_3P -metathesis barrier generating XY species is only ~ 5 kcal mol⁻¹, such that once radicals are generated an equilibrium would be expected to be established rapidly (see ESI† for details).

The DFT results suggested that Ar₂P* radicals (generated photolytically) were viable intermediates in the P,P-metathesis reactions and so this hypothesis was pursued experimentally.

Qualitative experimental kinetic studies of diphosphane metathesis

The aryl diphosphane metathesis reactions with A_2 – D_2 occurred readily (typically <1 h) under ambient conditions in common organic solvents (C_6D_6 , THF, PhMe, PhCl, CDCl₃ and CD_2Cl_2). By contrast, when the diphosphane E_2 , containing o-tolyl substituents, was one of the reaction partners, P,P-metathesis was initially only observed to take place in CDCl₃ (typically over 6–12 h) and CD_2Cl_2 (typically over more than 24 h); the anomalous behaviour observed for the P,P-metatheses involving E_2 will be discussed after consideration of the P,P-metatheses involving A_2 – D_2 .

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When cold (-78 °C) THF solutions of B₂ and C₂ were mixed and the P,P-metathesis reaction monitored by low-temperature ³¹P NMR spectroscopy, it was found that only traces of **BC** were detected at -80 °C after 1 h. By raising the temperature in increments of 20 °C, it was established that the P,P-metathesis at -20 °C progressed at a convenient rate to monitor the approach to P,P-metathesis equilibrium over a period of 2 h by ³¹P NMR spectroscopy (see Fig. 3). Although a detailed kinetic study was not attempted, the data do not appear to fit a reaction order of 1, 2 or 0.5 (see ESI†).

Evidence for radicals in the general P,P-metathesis reactions

In light of the computational results above suggesting that radicals may be involved in the mechanism of P,P-metathesis, experiments were carried out to determine if there was evidence of their presence. Two pieces of experimental evidence support the involvement of radicals in the P,P-metathesis process.

(1) When A_2 and C_2 were mixed in the presence of the radical scavengers 2,2,6,6-tetramethyl-1-piperidinyloxy (TEMPO) or tri(t-butyl)phenol (TTBP), the progress of the P,Pmetathesis reactions was greatly inhibited. Thus, a mixture of A2 and C2 proceeded smoothly to the metathesis equilibrium over a period of 2 h in CDCl₃, at -20 °C (with 35% conversion at 1 h); by contrast, when the same A₂/C₂ mixture was prepared in the presence of 5 equiv. of TEMPO, there was no formation of AC detected after 1 h. Moreover, when the reaction mixture containing TEMPO was allowed to warm to ambient temperature, <10% of AC was detected even after 16 h. Instead, in the presence of TEMPO, two singlet 31P resonances at +30.5 and +31.5 ppm were observed (ca. 10% of the total signal intensity), consistent with the formation of Ar₂P(O)-TEMPO (Ar = Ph, p-anisyl) adducts (Scheme 8) which would be expected to be formed by the interception of the phosphanyl radicals (Ar₂P*)

(2) When A₂ and B₂ were mixed in a standard NMR tube in THF at ambient temperatures, equilibration was essentially complete within ca. 15 min. However, when the same A₂/B₂ mixture in THF was prepared in an amberised NMR tube, to minimise the transmittance of UV and visible radiation, the

$$N-\dot{O} + Ar_2\dot{P}$$
TEMPO

Scheme 8

equilibration to the same degree took ca. 150 min. This is consistent with diphosphane photodissociation being a significant component of the mechanism of P,P-metathesis.

The UV-vis spectra of A_2 , C_2 and E_2 are shown in Fig. 4. Each spectrum has a major absorbance between 225 and 250 nm, and a shoulder at ca. 270 nm. TD-DFT calculations for A2 were carried out using a range of functionals (CAM-B3LYP, PBE0 and M06-2X) and good agreement with experiment was found with CAM-B3LYP. These calculations suggest that the major peak in each spectrum corresponds to coincident aromatic $\pi \to \pi^*$ and $S_0 \to S_2$ (see Fig. 5(b)) transitions and the shoulder peak corresponds to $S_0 \rightarrow S_1$ (see Fig. 5(a)). The transition electron density plots (Fig. 5) indicate that both transitions result in a significant delocalisation of electron density from the P lone pairs. The first triplet state, accessible from

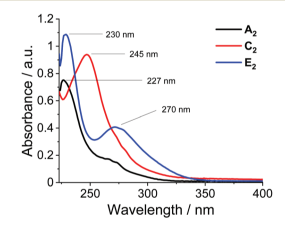


Fig. 4 UV-vis spectra of A_2 , C_2 and E_2 in CH_2Cl_2 (20 μ M).

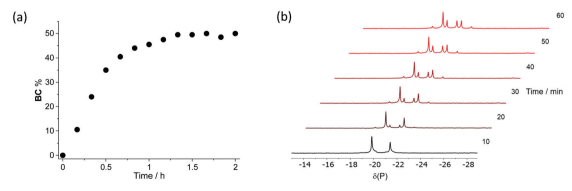


Fig. 3 (a) Plot of heterodiphosphane BC formation as a function of time over 2 h; (b) $^{31}P^{1}H$ NMR spectra of the reaction between B_2 and C_2 at -20 °C in THF, shown in 10 min increments over the course of 1 h (with inverse-gated decoupling). See Fig. 1 for the peak assignments.

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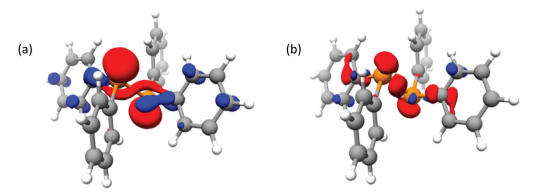


Fig. 5 Transition electron density difference plots with the CAM-B3LYP/def2-TZVPP//PBE0-D3BJ/def2-SVP level of theory. (a) $S_0 \rightarrow S_1$ and (b) $S_0 \rightarrow S_2$. Red and blue correspond to areas of depleted and excess electron density, respectively. Isosurface plotted at a contour of

either S₁ or S₂ via intersystem crossing, is dissociative, leading to spontaneous formation of phosphinyl radicals. There are small differences in the three spectra shown in Fig. 4. As expected, the p-OMe groups in C₂ produce a bathochromic shift in the $\pi \to \pi^*$ band and there is a long 'tail' that extends towards the visible region, which explains why C2 is pale yellow.

The evidence in support of radicals being present led to the radical chain mechanism proposal in Scheme 9 where homolysis of the Ar₂P-PAr₂ is followed by attack of the Ar₂P* on the second diphosphane Ar'2P-PAr'2. The termination steps are then the radical couplings with P-P bond formation.

The photodissociation requires UV radiation and it appears that there is sufficient background UVA-UVB radiation (280-400 nm) for the initiation step to proceed.

The two degenerate propagating steps, where Ar₂P* and Ar'₂P* react with their respective dimers Ar₂P-PAr₂ and Ar'₂P-PAr'₂, are omitted from Scheme 9. The reverse P,P-metathesis process

initiation PAr'2 propagation PAr₂ termination

Scheme 9

would also be initiated by photodissociation of Ar₂P-PAr'₂. The P,P-metathesis reactions generally proceed with a high degree of chemoselectivity but traces of Ar₂PH were often evident after several days of reaction which would arise from termination via abstraction of an H radical, presumably from the solvent.

The radical mechanism for diphosphane metathesis in Scheme 9 is reminiscent of the radical mechanism for disulfide metathesis (Scheme 10) arrived at from calculations by Ruipérez et al. 52 and from experimental observations by Asua et al.53 DFT and MD calculations were applied to two mechanisms of diaryldisulfide metathesis: a [2 + 2] concerted metathesis and a [2 + 1] radical chain reaction initiated by sulfanyl radicals (ArS*).47 The transition state in the concerted process was calculated to be too high in energy to be viable, whereas the radical process was viable provided the ArS' radicals could be accessed. This reasoning mirrors ours for the mechanisms proposed for diphosphane metatheses shown in Scheme 7.

It has been reported that the diaryldisulfide metathesis between diphenyldisulfide and di(4-aminophenyl)disulfide proceeds spontaneously to equilibrium in ca. 12 h; this was accelerated by UV photolysis, and TEMPO completely inhibited the equilibration.⁵³ These observations made on the disulfide metathesis reactions (and analogous diselenide studies)54 are very similar to our observations on the diphosphane metatheses.

Anomalous kinetics with (o-Tol)₂P-P(o-Tol)₂

The four P,P-metathesis reactions between tetra(o-tolyl)diphosphane (E_2) and A_2-D_2 did not appear to proceed at all in THF, PhMe, PhCl, PhF or C₆D₆ under ambient conditions after 72 h but do proceed in the chloroalkanes CH₂Cl₂ and CHCl₃, albeit slowly (see Table 1, entries 7-10). To probe this anomalous behaviour further, we have investigated the A₂/E₂ system (Fig. 6) in more detail including approaching the equilibrium starting from an A_2/E_2 mixture and starting from isolated AE.

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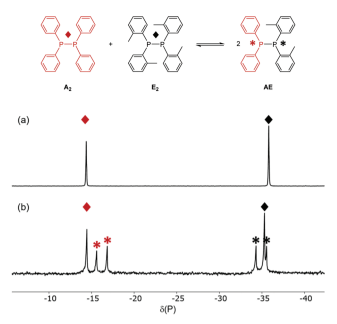


Fig. 6 $^{31}P\{^{1}H\}$ NMR spectra for the mixture of A_{2} and E_{2} : (a) in toluene after 48 h, showing no detectable P,P-metathesis; (b) in CDCl₃ after 6 h, showing equilibrium with AE has been established.

The progress of the A₂/E₂ metathesis in CDCl₃ was followed by 31P NMR spectroscopy (see ESI† for details) with a spectrum recorded every 10 min for 12 h (see Fig. 7). The approach to equilibrium was measured from integration of the 31P NMR signals to produce the plot shown in Fig. 7. Analysis of the curve revealed that it was not consistent with the integrated equations for an order of reaction of 0, $\frac{1}{2}$, 1 or 2; this indicates that the kinetics are complex, which would be expected if more than one mechanistic pathway was operating (see below).

Evidence for radicals in the anomalous (o-Tol)₂P-P(o-Tol)₂ metathesis reactions

There are several pieces of evidence supporting a role for radicals in the P,P-metathesis equilibrium shown in Fig. 6:

- (1) The lack of any reaction between A2 and E2 in THF after 72 h was replaced by 30% AE formation when the solution was irradiated with a UV lamp for 2 h. Similarly, no equilibration was observed in a THF solution of pure AE over a period of 72 h but proceeded smoothly over 2 h upon irradiation of the same solution with a UV lamp. This suggests that the photodissociation mechanism shown in Scheme 6 for the general P,P-metathesis (Table 1, entries 1-6) is viable for the A₂/E₂ metathesis but requires a more intense source of UV radiation than background UV for the initiation step.
- (2) The slow A₂/E₂ metathesis equilibration in CDCl₃ which takes place over a period of 12 h was extinguished in the presence of 4 equiv. of the radical scavenger TEMPO, as no AE was detected after 16 h.
- (3) Pulse-sonication (20 kHz) at 0 °C of a CHCl₃ or CH₂Cl₂ solution of A2 and E2 reduced the time to establish equilibrium from 12 h to <1 h. However, sonolysis had no accelerating effects on the reaction of A2 with E2 in THF or toluene. Fritze and von Delius reported the metathesis of disulfides (Scheme 10) in chloroform was accelerated by sonolysis and suggested that this proceeded via RS' radicals formed from the reaction of disulfides with Cl* generated by sonolytic chloroform degradation.16
- (4) The onset of equilibration of solutions of AE in CDCl₃ was preceded by an induction period of up to 6 h; moreover, the time required to establish the P,P-metathesis equilibrium from mixtures of A2/E2 in CDCl3 varied from 6 to 12 h. These observations of rate-inconsistency are typical of radical processes.55

In addition to the evidence for a radical-promoted process, it is shown below that there is evidence of chlorophosphine involvement in the A_2/E_2 metathesis.

Evidence for chlorophosphine intermediacy in the anomalous (o-Tol)₂P-P(o-Tol)₂ metathesis reactions

Under the conditions where a mixture of A₂ and E₂ in CDCl₃ would reach equilibrium with AE within 12 h, the presence of H_2O (100 equiv.) suppressed the reaction to the extent that no

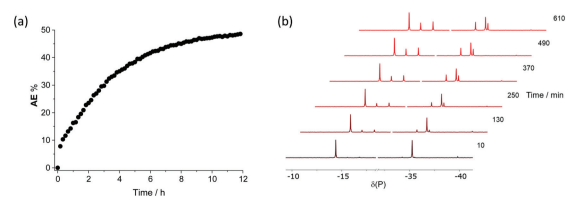


Fig. 7 (a) Plot of the data obtained from integration of the ³¹P(¹H) NMR signals for the AE formed in the reaction between diphosphanes A₂ and E₂; (b) $^{31}P^{1}H$ NMR spectra in CDCl₃ obtained by sampling over 12 h. See Fig. 6 for the peak assignments.

AE was detected even after 20 h (indicating that <1% **AE** had been formed). Instead, several minor signals (*ca.* 20% in total) were present and those at δ + 23.5, +21.4 were tentatively assigned to $Ar_2P(O)H$ (Ar = Ph or *o*-Tol). These observations are consistent with the water reacting rapidly with the Ar_2PCl compounds that are formed. It is notable that the addition of water to the P,P-metathesis reaction mixture of A_2 with C_2 in CDCl₃ had no discernible effect on the rapid formation of **AC** which indicates that Ar_2PCl is not involved in the P,P-metathesis for this typical system.

When a solution of A_2 alone in CHCl₃ was irradiated with UV, some conversion to Ph₂PCl was observed directly. The stoichiometric reaction of A_2 with $(o\text{-Tol})_2$ PCl in CHCl₃ immediately produced a mixture of A_2 , AE, and Ar₂PCl (Ar = Ph, o-Tol) (Scheme 11) along with several unidentified P-containing species. After 4 h, E_2 was detected along with small amounts of secondary phosphines Ar₂PH (Ar = Ph, o-Tol). These observations confirm that Ar₂PCl compounds are plausible promoters of P,P-metathesis.

To explore whether Ar_2PCl could catalyse P,P-metathesis, a sub-stoichiometric amount (5 mol%) of Ph_2PCl was added to a chloroform solution of pure **AE**. The equilibration (Scheme 11) was then monitored by $^{31}P\{^{1}H\}$ spectroscopy and compared with the same process carried out in the absence of Ph_2PCl . In the absence of Ph_2PCl , equilibrium was established in *ca*. 70 h, whereas in the presence of 5 mol% of Ph_2PCl , equilibrium was established in *ca*. 7 h (see ESI†).

The key observations on the P,P-metathesis reaction between E_2 and A_2 are that: (1) the solvent (CHCl₃ or CH₂Cl₂) is critical for the reaction to proceed under ambient conditions; (2) the reaction has the characteristics of being radical-initiated; (c) Ar_2 PCl catalyses the P,P-metathesis.

It is known that CHCl₃ is a source of Cl* radicals (especially under photolysis^{56–60} or sonolysis^{61–64}) and therefore the process shown in Scheme 12 is proposed in which the role of

Cl' is twofold: (a) to generate Ar_2PCl which promotes the P,P-metathesis process (Scheme 11); (b) to form Ar_2P^* and initiate a radical chain P,P-metathesis process similar to Scheme 9. DFT studies (see ESI†) show that initiation by attack of Cl' on Ar_2P-PAr_2 generating Ar_2P^* (Scheme 12) is feasible: $\Delta G = -35.3$ kcal mol^{-1} and $\Delta G^{\ddagger} = 3.4$ kcal mol^{-1} (from the $Ph_2P-PPh_2\cdots Cl^*$ association complex).

The slow rate of the P,P-metathesis reaction between the o-tolyl diphosphane \mathbf{E}_2 and \mathbf{A}_2 contrasts with the rapid rate of the P,P-metathesis reaction between the isomeric p-tolyl diphosphane \mathbf{B}_2 and \mathbf{A}_2 . The explanation for the lower rates therefore lies in the proximity of the o-CH $_3$ group to the P-reaction centre. We have proposed that photolytic Ph $_2$ P radical formation initiates the P,P-metathesis reactions of \mathbf{A}_2 (see above) and, since the UV spectrum of \mathbf{E}_2 resembles that of \mathbf{A}_2 (see Fig. 4), both Ph $_2$ P and (o-Tol) $_2$ P radicals should be similarly accessible. We propose two tentative explanations for the role of the o-CH $_3$ groups in \mathbf{E}_2 . (1) The bulky o-tolyl substituents may be shielding \mathbf{E}_2 from radical attack by Ph $_2$ P radicals or hindering radical attack by (o-Tol) $_2$ P on \mathbf{A}_2 . Some support for a steric effect of an ortho-alkyl group on the rate of radical-initiated P,P-metathesis comes from calculations of

Scheme 11 P,P-Metathesis promoted by Ar₂PCl.

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Scheme 13

 ΔG^{\dagger} for ${\rm Ar_2P^{\bullet}}$ attack on ${\rm Ar_2P-PAr_2}$ where ${\rm Ar}={\rm Ph}, o\text{-}{\rm C_6H_4Me}$ or $o\text{-}{\rm C_6H_4}^t{\rm Bu}$ which gave values of 5.5, 6.7 and 13.3 kcal ${\rm mol}^{-1}$ respectively (see ESI† for details). These show that, although the calculated ΔG^{\dagger} values for Ph and $o\text{-}{\rm C_6H_4Me}$ are similar, for $o\text{-}{\rm C_6H_4}^t{\rm Bu}$, the calculated value is significantly greater. (2) Alternatively, the $o\text{-}{\rm CH_3}$ groups may provide a means of quenching the radical chain by H* intramolecular abstraction from the proximal ${\rm CH_3}$ group to produce a benzylic radical by a process that is illustrated in Scheme 13. Calculations show that this would be highly unfavourable energetically (by 14.4 kcal ${\rm mol}^{-1}$) for the ground state $(o\text{-}{\rm Tol})_2{\rm P}^*$ radical but the initially formed P-centred diphosphane radical would have sufficient energy for H transfer to take place (see ESI† for details). The energetic C-centred radical could then abstract H* and the diphosphane reform thereby quenching the P,P-metathesis.

Both of these explanations are speculative and further work would be required to establish their veracity.

Conclusions

DFT calculations have shown that the thermodynamic stability of unsymmetrical diphosphanes (Z_2P-PZ_2), with respect to their symmetrical analogues (Z_2P-PZ_2 and Z'_2P-PZ_2), should increase with increasing difference in the electronegativity and/or steric bulk of the PZ_2 and PZ'_2 constituents. In these terms, the observed stability of $Me_2P-P(CF_3)_2$ and $^tBu_2P-PPh_2$ can be understood.

The unsymmetrical diphosphanes ($Ar_2P-PAr'_2$) have been shown to be in finely balanced equilibria with their symmetrical analogues Ar_2P-PAr_2 and $Ar'_2P-PAr'_2$ which reflects their insensitivity to the aryl substituents employed (Me, MeO, CF₃). Significantly, the P,P-metathesis equilibria with the tetra-aryl-diphosphanes were generally established rapidly (<10 min) at ambient temperatures, in a range of solvents, even in dilute solution; furthermore, the rate was increased by photolysis and inhibited by radical scavengers. The experimental and computational evidence points to the involvement of Ar_2P^* radicals generated photochemically.

Exceptional behaviour has been observed in the P,P-metathesis reactions involving $(o\text{-Tol})_2\text{P-P}(o\text{-Tol})_2$. These were only observed under UV-radiation or when the reactions were carried out in CHCl₃ and CH₂Cl₂. The evidence supporting the involvement of Ar₂PCl in the metatheses with $(o\text{-Tol})_2\text{P-P}(o\text{-Tol})_2$ in chlorinated solvents includes the effect of the presence of H₂O: no P,P-metathesis is observed and the formation of hydrolysis products Ar₂P(=O)H is detected. By contrast, the P,P-metatheses involving other Ar₂P-PAr₂ proceed uninhibited by the presence of water and no Ar₂P(=O)H species are detected.

The facility with which diphosphanes undergo P,P-metathesis is similar to the S,S-metathesis with disulfides and Se, Se-metathesis with diselenides. This augurs well for the application of diphosphanes in dynamic covalent chemistry.

Conflicts of interest

There are no conflicts to declare.

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