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Reactivities of allenic and olefinic Michael acceptors towards phosphines[†]

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The kinetics of the reactions of tributylphosphine with allenic and olefinic Michael acceptors in dichloromethane at 20 °C was followed by photometric and NMR spectroscopic methods. Combination with DFT-calculated methyl anion affinities revealed the relevance of retroaddition barriers in phosphine-catalysed reactions when mixtures of allenic and olefinic substrates are used.

In 1995, the Lu group discovered (3+2)-cycloadditions, in which Bu_3P or Ph_3P are used as Lewis-basic catalysts to furnish cyclopentenes from mixtures of alkyl allenoates and electron-deficient alkenes (Fig. 1).¹ In recent years, the versatility of Lu cycloadditions was expanded by the development of several catalytic asymmetric versions, which were also applied to the synthesis of core units of natural products.² Computational and kinetic studies showed that phosphonium–dienolate formation can be considered to be the rate-determining step in the catalytic cycle.^{2a,3}

The rates of adduct formations, in which only one new bond is formed between an electrophile and a nucleophile, can be discussed in a broader context and beyond the limitations of structurally analogous compound classes when the Mayr–Patz eqn (1) is used.^{4–7} Thus, the nucleophilic reactivity of R₃P catalysts in a certain solvent is described by the two parameters N (nucleophilicity) and s_N (susceptibility) in eqn (1).⁸ The reactivities of Michael acceptors are characterised by electrophilicity parameters *E*. It has been shown that once *E*, *N* and s_N of reaction partners in polar reactions are known the secondorder rate constant k_2 for adduct formation can be predicted with an accuracy within two orders of magnitude.^{5,6}

$$\log k_2(20 \ ^\circ\mathrm{C}) = s_{\mathrm{N}}(N+E) \tag{1}$$

Many of the R_3P -catalysed reactions are performed with mixtures of two competing electrophiles, which are often combined in a way that one of the electrophiles carries an spand the other one an sp²-hybridised carbon as the electrophilic centre.^{1,2,9} In this work, we set out to determine the kinetics of the adduct formation of R_3P with electron-deficient olefins and a set of alkyl and phenyl allenoates to gain a deeper understanding of the factors that influence the initial step of the related R_3P -catalysed organic reactions.^{1,2,9}

First, we characterised the vinyl phosphonium triflates 3 obtained by Ph_3P reactions with **1a**, **1d**, **1f**, and **1i** (Fig. 2A) by spectroscopic methods (ESI[†]). It is reasonable to assume that the entire set of $R_3P + 1a-1j$ reactions that we followed kinetically also yield vinyl phosphonium triflates 3. The kinetics of the carbon-phosphorus bond-formations between R_3P and **1a-1j** was followed by spectroscopic methods.



Fig. 1 Mechanism of the phosphine-catalysed Lu cycloaddition (E = ester group, Acc = electron-accepting group).

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Fig. 2 (A) R_3P additions to **1** and (B) reference reaction for the determination of methyl anion affinities (MAA) by quantum-chemical methods.

Reactions of Bu₃P with alkyl allenoates 1a-1i in CH₂Cl₂ at 20 °C were monitored photometrically by following the decrease of the UV absorptions of 1 at or close to their absorption maximum (λ_{max} = 242–262 nm). The Bu₃P addition reactions to 1 give rise to polymerisation, which can be avoided by trapping the zwitterionic adducts 2 with tributylphosphonium triflate (TBPT) as the proton source ($[TBPT]/[1]_0 = 2-3$). TBPT is a weak Brønsted acid in CH₂Cl₂ and does not activate the electrophiles 1 by hydrogen-bonding to an oxygen of the ester group (as shown for the combination of 1f + TBPT, ESI,† Fig. S7). To simplify the kinetic evaluation of the second-order reactions, we used the Bu₃P in at least 10-fold excess relative to the initial electrophile concentrations $[1]_0$. Hence, the decrease of absorptions A of 1 could be fitted by the mono-exponential decay function $A = A_0 \exp(-k_{obs}t) + C$ to determine the (pseudo) first-order rate constants k_{obs} (s⁻¹).

For each Bu₃P + **1** pair, k_{obs} was determined at four or five different Bu₃P concentrations, which made it possible to calculate the second-order rate constants k_2 (M⁻¹ s⁻¹) from the slope of the linear correlation of k_{obs} with [Bu₃P]₀. Fig. 3 visualises this procedure for the Bu₃P addition to **1f**.

Rate constants of the reactions of Ph_3P with **1a–1j** were determined by ¹H NMR spectroscopy using mesitylene as an internal integration standard and collidinium triflate (CT) as a proton source ([CT]/[**1**]₀ = 2). Generally, Ph_3P additions to **1** are endergonic with retroadditions being faster than the addition reactions. Trapping of the zwitterionic intermediates **2** by CT is thus necessary to observe the kinetics of the addition (forward)



Fig. 3 (A) Kinetics of the reaction of Bu_3P with **1f**: mono-exponential decay of the absorbance A in the reaction of Bu_3P (12.3 mM) with **1f** (0.121 mM); (B) linear correlation of observed rate constants k_{obs} with $[Bu_3P]_0$.

Table 1 Second-order rate constants (k_2 , in M⁻¹ s⁻¹) for the additions of Bu₃P and Ph₃P to phenyl or alkyl allenoates **1** in dichloromethane at 20 °C

Electrophiles 1	$k_2(\mathrm{Bu}_3\mathrm{P})^a$	$k_2(Ph_3P)^b$	MAA ^c	\mathbf{PA}^d
1a	5.52	$1.19 imes10^{-1}$	182.5^{e}	-15.0
1b	1.16	$1.70 imes10^{-2}$	163.1^{e}	-22.6
1c	1.10	$1.90 imes10^{-2}$	165.5	-23.7
1d	$9.55 imes10^{-1}$	$2.33 imes10^{-2}$	171.2	-15.9
1e	8.40×10^{-1}	$1.10 imes10^{-2}$	167.6^{e}	-26.5
1f	$6.35 imes10^{-1}$	$7.67 imes10^{-3}$	163.4^{e}	-29.8
1g	5.00×10^{-1}	8.05×10^{-3}	152.7	-29.5
1ĥ	2.47×10^{-1}	2.39×10^{-3}	153.9	-33.3
1i	2.01×10^{-1}	4.59×10^{-3}	142.5	-26.0
1j	1.96×10^{-2b}	3.49×10^{-4}	133.6	-38.3

^{*a*} Photometry, CH₂Cl₂, 20 °C. ^{*b*} ¹H NMR spectroscopy, CD₂Cl₂, 20 °C. ^{*c*} MAA as defined in Fig. 2B [in kJ mol⁻¹ at SMD(DMSO)/B3LYP/6-311++G(3df,2pd)//B3LYP/6-31G(d,p) level of theory, with Truhlar quasi-harmonic treatment]. ^{*d*} Phosphine affinities, PA, as defined in ESI, Table S42 [in kJ mol⁻¹ at PCM(DCM,ua0)/B3LYP-D3/6-31+G(d,p) level of theory, with Truhlar quasi-harmonic treatment]. ^{*e*} MAA values from ref. 10.

reaction. By obeying the conditions for (pseudo)first-order kinetics, that is $[Ph_3P]_0/[1]_0 > 9$, the time-dependent decrease of [1] could be fitted by the mono-exponential decay function to yield the rate constants k_{obs} . NMR experiments at four different $[Ph_3P]_0$ made it possible to determine the second-order rate constants k_2 for the Ph₃P additions to the electrophiles **1a–1j** from the slope of the linear correlation of k_{obs} with $[Ph_3P]_0$. Rate constants (k_2) for the reactions of Bu₃P with **1j** were determined analogously. Data of the individual kinetic measurements for the nucleophilic attack of Bu₃P and Ph₃P at the allenoates **1** are given in the ESI.† The experimentally determined second-order rate constants k_2 are compiled in Table **1**.

The Bu₃P-based reactivity scale (log k_2) for **1a–1j** is depicted in Fig. 4A. The phenyl ester **1a** is the strongest electrophile of the studied allenoates without further substituents at the cumulated π -system and reacts about 5 to 9 times faster with Bu₃P than the analogous benzyl (**1b**), methyl (**1e**), or ethyl esters (**1f**).¹¹ Replacement of the ethyl by a *t*-butyl group (**1f** \rightarrow **1h**) attenuates reactivity by a factor of 3. An additional methyl



Fig. 4 (A) Relative reactivities of **1a–1j** in reactions with Bu_3P (in CH_2Cl_2 , 20 °C). (B) Different stereoelectronic effects in **1d** and the open-chain analogue **1j**.

group (**1g**) at the 4-position of the alkyl allenoate has almost no influence on the reactivity of the electrophilic π -system, which remains at the level of the γ -unsubstituted **1f**. Extending the π -system by a terminal phenyl group enhances the reactivity of **1c** only slightly if compared to that of **1f**, probably because the phenylvinyl moiety in **1c** is perpendicular to the reactive π -system. In contrast, substituents in 2-position of **1** have a greater effect, and Bu₃P reacts slower by a factor of 32 with **1j** than with the parent **1f**. This reactivity gap can be reduced by attaching an electron-accepting group to the C-2 substituent. Thus, **1i** reacts 10-fold faster with Bu₃P than **1j** and is only 3 times less electrophilic (towards Bu₃P) than **1f**.

Consistent with previous work on the relative reactivity of open chain esters and lactones,¹² α -allenic γ -butyrolactone **1d** is a significantly more reactive electrophile than **1j**. The increase in electrophilic reactivity is explained by the different conformations of the CO–OR bonds, which is preferably in the *s*-(*Z*) conformation in **1j** but fixed in the unfavourable *s*-(*E*) conformation in **1d** (Fig. 4B).^{12,13} Ineffective $n_O \rightarrow \sigma^*_{CO}$ interactions enhance the electron-deficiency of the reacting π -system, which enables the versatile use of allenic lactones in organic synthesis.¹⁴

Reactivity of **1** towards Bu_3P is transferrable to other R_3P as shown by the linear correlation ($r^2 = 0.9636$, n = 10) with a slope (1.02) close to unity for the Ph₃P vs. Bu₃P comparison (Fig. 5).

To gain a better understanding of R_3P -catalysed Lu reactions it is crucial to compare the R_3P reactivities of **1** with those of competing electrophiles, which are typically olefinic Michael acceptors. It was previously shown that MAAs of olefinic Michael acceptors correlate linearly with their Mayr *E* parameters.¹⁵ MAA values have also been applied to rationalise R_3P -catalysed (3+2) annulations of **1a**, **1b**, **1e**, and **1f** with 2-aminoacrylates.¹⁰ The data in Table 1 now show that the DFT-calculated MAA values for **1** (Table 1) are linearly related with their electrophilic reactivities towards the investigated R_3P nucleophiles, that is, Bu_3P and Ph_3P (ESI,† Fig. S1 and S2). A linear correlation of similar quality was obtained when $\log k_2$ for **1** + Ph_3P reactions were plotted against phosphine affinities, PA, which are defined analogously to MAA but use Ph_3P instead of the methyl anion as the Lewis base (Fig. S3, ESI†).

The nucleophilic reactivity of Bu₃P has previously been characterised by N = 15.49 ($s_{\rm N} = 0.69$) on the basis of the kinetics of its additions to benzhydrylium ions (E > -10.04).¹⁶ To avoid



Fig. 5 Linear relation of $\log k_2(Bu_3P)$ and $\log k_2(Ph_3P)$ for reactions with **1**.



Fig. 6 Linear correlation of $\log k_2(\operatorname{Bu}_3 P)$ with the Mayr *E* parameters of structurally diverse Michael acceptors **4a–o** and benzhydrylium ions **4p–r** (k_2 in dichloromethane at 20 °C, see ESI† for details of the kinetic measurements, Table S1 (ESI†) gathers the data used to construct the correlation).

long range extrapolations for predicting reaction rates with Michael acceptors, we determined the kinetics of further reactions of Bu₃P with the structurally diverse neutral and positively charged electrophiles **4a–4r** of known Mayr *E*.⁷ Second-order rate constants k_2 (CH₂Cl₂, 20 °C) were determined by applying the photometric methods described above for the **1** + Bu₃P reactions (see ESI† for individual rate constants). Fig. 6 shows that log k_2 for the additions of Bu₃P to **4a–4r** follow a linear correlation ($r^2 = 0.9635$, n = 18) over a range of 16 units on the Mayr *E* scale, which gives $N(Bu_3P) = 19.11$ and $s_N = 0.48$. Applying the k_2 values for reactions of **1** (Table 1) in the correlation depicted in Fig. 6 indicates that **1a–1j** are located in the reactivity range -22.7 < E < -17.5.

In classical Lu reactions, the R_3P catalyst first attacks the alkyl allenoate. The catalytic cycle continues with the reaction of a C-nucleophilic zwitterion with the second Michael acceptor in the reaction mixture. If the kinetics of the R_3P reactions with the two competing electrophiles would be the decisive factor, Lu reactions could be expected to occur only if 1 is more reactive towards the R_3P catalyst than the competing Michael acceptor (*e.g.*, 4). This is not always the case, however. Already in the first publication on the R_3P -catalysed cycloaddition, the Lu group¹ used electrophiles with Mayr E < -16.8, that is, with comparable or even slightly higher electrophilicity than for 1. This indicates that the thermodynamics for R_3P adduct formation is another crucial factor for the success of Lu reactions.

For reactions of vinyl cations with nucleophiles it has been observed that sp/sp^2 rehybridisation occurs *via* higher Marcus intrinsic barriers than for reactions that involve sp^2/sp^3

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Fig. 7 (A) Separate linear relationships between the electrophilicities *E* of acceptor-substituted olefins **M1–M44** and allenes **1** with the calculated methyl anion affinities (MAA) (molecular structures of **M1–M44** and data used to construct the linear correlation are given in ESI,† Table S2; electrophilicities *E* for allenoates **1** estimated by applying the $k_2(Bu_3P)$ values from Table 1 in the correlation equation for Bu_3P additions to Michael acceptors in Fig. 6). (B) MAA values and electrophilicities *E* for Michael acceptors **4s–4v**.

rehybridisations.^{17,18} Analogously, in this work the higher MAA for 1 than for equally reactive olefinic Michael acceptors^{15a} (Fig. 7) along with the experimental kinetic data indicate that the main difference between 1 and Michael acceptors, such as 4, are the Marcus intrinsic barriers for the R_3P addition at the differently hybridised electrophilic centres.

Even if the rate constants for the electrophile/R₃P additions are of the same order of magnitude, the MAA values show that allenoates 1 are considerably stronger Lewis acids than olefinic Michael acceptors or imines, such as 4t. As a consequence, the differences in the barriers for the retroadditions differentiate the two competing classes of electrophiles. Owing to their higher energetic barrier for retroaddition, only the allenic electrophiles 1 generate sufficiently high concentrations of reactive zwitterions, which are the pivotal intermediates for the subsequent ring-forming reactions. Thus, allenoates 1 are capable to compete with much stronger electrophiles. For example, 1f (MAA = 163 kJ mol⁻¹) can be used as a partner for the more electrophilic yet less Lewis acidic 4s, 4t, or 4u in Lu cycloadditions.¹⁹⁻²¹ Benzylidenemalononitrile (4v) seems to be an exception. However, despite of its high MAA, 4v reacts reversibly with Bu₃P and does not form an adduct with Ph₃P (ESI⁺). Yet, free 4v traps efficiently the zwitterion generated by Ph₃P and **1f** to yield cyclopentenes.^{22,23}

The reactivities of allenic and olefinic Michael acceptors have been calibrated towards P-nucleophiles through determining the kinetics of their reactions with Bu_3P . Allenoates **1** are weaker electrofuges as well as weaker electrophiles than Michael acceptors **4** of similar Lewis acidity because of the higher intrinsic barriers for sp/sp^2 rehybridisation. The kinetic and thermodynamic data in this work will be instrumental for the design of novel R_3P -catalysed reactions with alkyl allenoates.

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Conflicts of interest

There are no conflicts to declare.

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