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Diphosphination of CO<sub>2</sub> and CS<sub>2</sub> mediated by frustrated Lewis pairs - catalytic route to phosphanyl derivatives of formic and dithioformic acid

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The first example of CO<sub>2</sub> diphosphination is described. Reactions of unsymmetrical diphosphanes with CE<sub>2</sub> (E = O, S) catalyzed by BPh<sub>3</sub> insert a CE<sub>2</sub> molecule into the P-P bond with formation of the products of the general formula R<sub>2</sub>P-E-C(=E)-PR<sub>2</sub>. The obtained CO<sub>2</sub> adducts arise from synergistic interaction of diphosphane and borane with CO<sub>2</sub> molecule.

Carbon dioxide is an inexpensive, safe and abundant building block for the production of chemicals and fuels. Studies on the direct activation and transformation of carbon dioxide with simple nonmetallic systems have become one of the most active branches of modern catalysis research. Amid the great variety of compounds capable of activating CO<sub>2</sub>, the frustrated Lewis pairs (FLPs)<sup>1</sup> reported by Stephan, which are combinations of sterically hindered Lewis acids and bases that do not form classical adducts, contributed significantly to this field.<sup>2-6</sup> In the case of FLP systems based on phosphorus as the basic center, the activation of CO<sub>2</sub> proceeds via the formation of a P-C bond, and depending on the type of reactive acidic site, B-O,<sup>6,7</sup> AI-O<sup>8,9</sup> or Ga-O<sup>10</sup> bonds often in a reversible manner. An interesting feature of FLP-activated  $CO_2$  is that it can be converted into methanol under mild conditions.<sup>8,11</sup> Apart from phosphines that usually act as P-containing Lewis bases, there are also reports on P-P bond-containing systems such as symmetrical diphosphanes that when used with an equimolar amount of highly Lewis acidic  $B(C_6F_5)_3$  were found to react with  $H_2$  and phenylacetylene.<sup>12,13</sup> Although they are expensive,  $B(C_6F_5)_3$  and its fluorinated derivatives are widely employed in FLPs due to their high reactivity towards small molecules. Notably, the presence of a Lewis acidic component is not necessary for capturing CO2. Recently, it was reported that highly nucleophilic phosphines can reversibly form simple adducts with CO2.14 Despite wide laboratory and industrial

5 mol% BPh<sub>3</sub>

CO

5 mol% BPh<sub>3</sub>

iPr<sub>2</sub>

Et<sub>2</sub>

Et<sub>2</sub>

applications, the fixation of CS2 is not well studied. To date,

there are only a few reports regarding the activation of CS<sub>2</sub> by

FLPs involving the formation of Lewis base-acid adducts LB-C(S)-S-LA<sup>7,15,16</sup> and the diphosphination of  $CS_2$  with persistent

 $(H_2C)_2(NDipp)_2P \bullet$  radicals generated in solution from the parent symmetrical diphosphane.<sup>17</sup> Unsymmetrical diphosphanes

were also utilized in diphosphination reactions; however, they

were reacted with activated acetylenes.18,19 To the best of our

knowledge, there are no reports on the activation of small

Based on our long-term interest in the chemistry of compounds

possessing P-P functionalities<sup>20–27</sup> and the recent developments

in FLP chemistry, 10,28 we decided to study the reactivity of

unsymmetrical diphosphanes as Lewis basic components in

combination with weakly Lewis acidic BPh3. From the set of

recently developed systems with polarized P-P bonds,<sup>29</sup> we

selected three species, tBu<sub>2</sub>P-P(NiPr<sub>2</sub>)<sub>2</sub> (1), tBu<sub>2</sub>P-P(NiPr<sub>2</sub>)(NEt<sub>2</sub>)

(2) and  $tBu_2P-P(NEt_2)_2$  (3), with highly nucleophilic P-atoms. We

inorganic molecules, such as CO<sub>2</sub> or CS<sub>2</sub>,

unsymmetrical systems with polarized P-P bonds.

involving

Scheme 1. Reactions of 1-3 with CO2 catalyzed by  $\mathsf{BPh}_3$ 

tBu

tBu

tBu

*i*Pr<sub>2</sub>N

Et<sub>2</sub>

3

tBu 3a

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found that mixtures of BPh<sub>3</sub> and a base (1, 2 or 3) not only form stable FLPs that do not quench each other but also react with  $CO_2$  and/or  $CS_2$  in an unusual manner. The reaction of a stochiometric mixture of 1 and BPh<sub>3</sub> in toluene under 1 atm of CO<sub>2</sub> at ambient temperature for 24 hours led to complete conversion of 1 into 1a (Scheme 1). tBu *t*Bu **1a** 1 5 mol% BPh3 7Ru *i*Pr<sub>2</sub>N Et<sub>2</sub>N tBu CO Et<sub>2</sub>N 2 *t*Bu **2a** 

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The <sup>31</sup>P NMR spectrum of the reaction mixture showed two doublets at 113.4 ppm (P(NiPr<sub>2</sub>)<sub>2</sub>) and 47.4 ppm (PtBu<sub>2</sub>) (1a). Relative to the signals of starting diphosphane 1, the doublet attributed to the P(NiPr<sub>2</sub>)<sub>2</sub> group is shifted downfield, whereas the signal of PtBu<sub>2</sub> is shifted upfield (for 1: 88.2 ppm and 62.6 ppm, respectively). Furthermore, the absolute value of the P-P coupling constant is significantly reduced from 358.2 Hz (1) to 14.5 Hz (1a), indicating the absence of a direct P-P bond in newly formed 1a. Surprisingly, the <sup>11</sup>B NMR spectrum of the reaction mixture only showed a signal attributable to BPh<sub>3</sub>. We did not observe any resonances for zwitterionic species bearing B-O moieties, which are typically formed in reactions of phosphorus-boron FLPs with CO2. This observation suggested the Lewis acid was serving as a catalyst in the described reaction. To verify this hypothesis, we performed reactions of 1 with  $CO_2$  in the presence of 25 mol% and 5 mol% BPh<sub>3</sub> and without the Lewis acid under the same conditions. Indeed, in the case of reactions involving catalytic amounts of BPh<sub>3</sub>, we observed complete conversion of 1 into 1a, while the absence of BPh<sub>3</sub> resulted in no reaction of 1 with CO<sub>2</sub>. Compound 1a was isolated by crystallization from the concentrated reaction mixtures at -20°C. The reaction with 5 mol% catalyst loading gave the highest isolated yield (90%) and purity as the isolated product was not contaminated by BPh<sub>3</sub>. Further spectroscopic analysis of isolated 1a also indicated the incorporation of a CO<sub>2</sub> molecule between the two phosphorous atoms. The <sup>13</sup>C NMR spectrum of 1a showed a very characteristic doublet of doublets at 178.2 ppm ( ${}^{1}J_{PC}$  = 29.4 Hz,  ${}^{2}J_{PC}$  = 4.4 Hz) attributed to the CO<sub>2</sub> moiety. Moreover, the IR data for 1a showed an absorption band at 1667 cm<sup>-1</sup>, which is typical for a C=O group. The X-ray diffraction data of 1a conclusively confirmed the insertion of CO<sub>2</sub> into the P-P bond (Figure 1). The CO<sub>2</sub> moiety is located between two phosphanyl groups; the PtBu<sub>2</sub> group is bound to the carbonyl C9-atom, while the P(NiPr<sub>2</sub>)<sub>2</sub> group is connected to one of the oxygen atoms. The C9-O1 (1.348(3) Å) and C9=O2 (1.205(3) Å) bond lengths are in the range typical for carboxylic esters (~1.33 - 1.41 Å and ~1.19 - 1.20 Å, respectively).<sup>30</sup> Furthermore, the P1-C9 (1.870(2) Å) and P2-O1 (1.736(2) Å) bonds are slightly longer than typical P-C and P-O single bonds (~1.80-1.85 Å and ~1.56-1.69 Å, respectively).30



Figure 1. X-ray structure of **1a** showing the atom-numbering scheme. The H atoms are omitted for clarity.

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As expected, the C9 atom exhibits trigonal planar/geometry ( $\Sigma$ 359.96°), while P1 and P2 are pyramidal 1213324.224°Cand12 302.94°, respectively). P1 and P2 are located almost in the same plane as the CO<sub>2</sub> moiety (torsion O2-C9-O1-P2: 5.9°). The geometries around N1 and N2 are almost planar due to the interaction of their lone pairs with the P-atoms, and the same structural feature was observed for 1.29 According to NBO analysis of structure 1a, the unusual planar array of P1-C9(=O2)-O1-P2 atoms results from sp<sup>2</sup> hybridization of both, C9 and O1 atoms. Furthermore, unhybridized p orbital of O1 (the one possessing a lone pair) interacts with antibonding  $\pi^*(C9-O2)$ orbital what is in accord with decreased occupancy (1.77) of the first one, and the increased occupancy (0.24) of the latter one. The colorless crystals of 1a are stable under an argon atmosphere or under vacuum, and regeneration of 1 was not observed. Moreover, warming the toluene solution of 1a to 50°C in the presence of BPh<sub>3</sub> for several days did not lead to decomposition of 1a. Interestingly, in contrast to 1, 1a is airstable both in the solid-state and in solution.

Experiments involving **2** and **3** indicated that small differences in the structural features of **1**, **2** and **3** have dramatic effects on the reactivity of these systems towards CO<sub>2</sub>. Stirring toluene solutions of **2** and BPh<sub>3</sub> (100 mol%/25 mol%/5 mol%) under a CO<sub>2</sub> atmosphere (1 atm) for 24 hours resulted in complete conversion of **2** into **2a** (Scheme 1). The spectral data of **2a** correspond to the spectral parameters determined for **1a** and confirm the insertion of a CO<sub>2</sub> molecule into the P-P bond. Unlike the reaction of **1**, the reaction of **2** with CO<sub>2</sub> is reversible. Heating a solution of **2a** with 5 mol% BPh<sub>3</sub> to 45°C for 24 hours under argon led to regeneration of **2** (molar ratio **2:2a**  $\approx$  4:1) (Figure 2). In contrast to **1** and **2**, **3** does not react with CO<sub>2</sub> under identical conditions, and corresponding adduct **3a** is not formed (Scheme 1).



Figure 2. Variable-temperature <sup>31</sup>P{<sup>1</sup>H} NMR spectra of **2a**.

To our delight, unsymmetrical diphosphane **1** reacted smoothly with  $CS_2$  in the presence of BPh<sub>3</sub>. The desired product **1b** was obtained upon mixing a toluene solution of **1** and BPh<sub>3</sub> (100 mol%/25 mol%/5 mol%) with an approximately threefold excess of  $CS_2$  at room temperature (Scheme 2). Monitoring the deep green reaction mixtures by  ${}^{31}P{}^{1}H{}$  NMR revealed complete conversion of **1** into **1b** after 48 hours. Diphosphanes

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 ${\bf 2}$  and  ${\bf 3}$  react with  $CS_2$  even in absence of BPh\_3, and the reactions are completed after 30 minutes. However, the formation of the  $CS_2$  adducts  ${\bf 2b}$  and  ${\bf 3b}$  proceeds by using sixfold excess of  $CS_2.$ 



Scheme 2. Reactions of 1-3 with  $CS_2$ .

The obtained products were characterized by  $^1\text{H},~^{31}\text{P}\{^1\text{H}\},$  and <sup>13</sup>C NMR, IR, and UV-Vis spectroscopy (see the ESI for details). Representative compound 1b exhibits two doublets at 106.6 ppm (P(N*i*Pr<sub>2</sub>)<sub>2</sub>) and 75.0 ppm (P*t*Bu<sub>2</sub>) ( ${}^{3}J_{PP}$  = 14.5) in its  ${}^{31}P{}^{1}H$ NMR spectrum, a signal of the  $CS_2$  moiety at 250.8 ppm (dd,  ${}^1J_{PC}$ = 64.5 Hz,  ${}^{2}J_{PC}$  = 20.9 Hz) in its  ${}^{13}C$  NMR spectrum and a C=S absorption band at 1045 cm<sup>-1</sup> in its IR spectrum. Collectively, these data confirm the incorporation of CS<sub>2</sub> molecules into the diphosphane. Compounds 1b and 2b were characterized by Xray analyses (Figure 3, Figure S3). The structural analysis of 1b and 2b showed that the CS<sub>2</sub> molecule was inserted in the P-P bond of the starting diphosphane with the formation of new P-C and P-S bonds. The geometries of 1b and 2b resemble that of 1a with the two phosphanyl groups linked by the CS<sub>2</sub> unit. Similar to 1a, the C9 atoms in 1b and 2b are in trigonal planar geometries, whereas the P-atoms are pyramidal. In contrast to 1a, the products of the reactions with CS<sub>2</sub> are air-sensitive. Moreover, **3b** is not stable in toluene solutions, and according to NMR studies, it rearranges into more thermodynamically stable species.

The reaction products of unsymmetrical diphosphanes **1-3** with  $CO_2$  or  $CS_2$  may be regarded as phosphanyl derivatives of formic and dithioformic acid. Very recently, free phosphinoformic acid was detected in the gas phase,<sup>31</sup> and its isolable derivatives are rare.<sup>32–34</sup> To the best of our knowledge, compounds with the R<sub>2</sub>P-E-C(=E)-PR<sub>2</sub> skeleton have not been described in the literature for E = O and are quite rare for E = S.<sup>17</sup>



Figure 3. X-ray structure of 1b showing the atom-numbering scheme. The H atoms are omitted for clarity

The presence of P,O-donor atoms in **1a** prompted us to examine its reactivity towards transition metal centers. The reaction of



1a with [(COD)PtMe<sub>2</sub>] in toluene at 50°C cleanly\_afforded 4

Scheme 3. Synthesis of 4.

(Scheme 3).

Slow evaporation of the reaction mixture led to precipitation of crystalline **4** (yield 90%), which was further characterized by <sup>1</sup>H, <sup>31</sup>P{<sup>1</sup>H}, and <sup>13</sup>C NMR spectroscopy and X-ray analysis. These data revealed that **1a** acts as a bidentate P-donor ligand, resulting in an almost planar five-membered metallacycle (Figure S4). Interestingly, the ligand geometry changes upon complexation. In free-**1a**, the P-atoms are *trans* to one another along the C–O bond, while in complexed-**1a**, they are *cis*.

Differences in the reactivity of diphosphanes 1-3 and the stability of the obtained  $CE_2$  adducts (E = O or S) were investigated by DFT calculations (see the ESI for details). The free energies  $\Delta G_{298}$  determined for the formation of the respective adducts confirmed that only 1a, 1b, and 2b should form stable products in exergonic reactions (Table 1), and the formations of 2a, 3a and 3b are not thermodynamically favorable. For the CS<sub>2</sub> and CO<sub>2</sub> adducts of **3**, only **3b** may be obtained in an endogenic reaction as an intermediate that rearranges to other presumably more stable species. By calculating the condensed nucleophilic Fukui functions (Table S12), we found that in each case, PtBu<sub>2</sub>, as the more nucleophilic center, preferred to attack electrophilic  $CE_2$ . Second, the less nucleophilic P-center reacts with electron-rich E-atoms. In general, we can assume that the more nucleophilic P-centers react more quickly. In contrast, the more nucleophilic phosphorus centers, the less stable products are formed.

Table 1. Values of enthalpies ( $\Delta$ H), free energies ( $\Delta$ G) of reactions **1a-3a** and **1b-3b** 

Reaction	1a	2a	3a	1b	2b	3b
<b>ΔH</b> kcal.mol-1	-13.0	-7.2	-5.6	-12.5	-8.4	-5.0
<b>ΔG</b> kcal.mol-1	-4.6	1.7	3.4	-4.1	-0.1	5.8

Experimental data provides us information about the reaction mechanism. We evidenced that presence of BPh<sub>3</sub> is crucial for obtaining **1a** and **2a**, and because of steric hindrance, borane does not form adducts with diphosphanes **1-3**. It is worth mentioning that neither BPh<sub>3</sub> nor diphosphanes **1-3** react with CO<sub>2</sub> alone. Even at low temperatures (between +25°C and - 50°C), we did not observe formation of simple CO<sub>2</sub>-adducts similar to those reported by Dielmann.<sup>14</sup> We also did not notice formation of any other reaction products. Hence, we may eliminate the radical mediated reaction mechanism as phosphanyl R<sub>2</sub>P radicals tend to react with the solvent or the

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other radical giving symmetrical diphosphanes. Altogether, these observations suggest that formation of **1a** and **2a** can be considered as a result of synergistic interactions of diphosphane (Lewis base) and borane (Lewis acid) with  $CO_2$  which is a characteristic reactivity pattern for FLPs.

Reactivity of diphosphanes **1-3** towards  $CS_2$  seems to be more diversified. Although presence of BPh<sub>3</sub> also facilitates the formation of respective  $CS_2$  adducts, in the case of highly nucleophilic systems **2** and **3**, reaction may proceed without Lewis acid by applying great excess of  $CS_2$ . Detailed reaction mechanisms are the subject of ongoing studies.

In conclusion, the reactivity of unsymmetrical diphosphanes towards  $CO_2$  and  $CS_2$  constitute a bridge between the reactivity of FLPs, highly nucleophilic phosphines and diphosphorus species. The diphosphination reactions described herein open new pathways for the activation of small molecules.

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## **Conflicts of interest**

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There are no conflicts to declare.

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