Article

Metal-Free Phosphination and Continued Functionalization of Pyridine: A Theoretical Study

Pan Du 1, Yuhao Yin 2, Dai Shi 2, Kexin Mao 2, Qianyuan Yu 2 and Jiyang Zhao 2,*

1 School of Life Science and Chemistry, Jiangsu Second Normal University, Nanjing 210013, China
2 School of Environmental Science, Nanjing Xiaozhuang University, Nanjing 211171, China
* Correspondence: jyzhao1981@163.com

Abstract: This study investigates the mechanism of metal-free pyridine phosphination with P(ΟEt)3, PPh3, and PAr2CF3 using density functional theory calculations. The results show that the reaction mechanism and rate-determining step vary depending on the phosphine and additive used. For example, phosphination of pyridine with P(ΟEt)3 occurs in five stages, and ethyl abstraction is the rate-determining step. Meanwhile, 2-Ph-pyridine phosphination with PPh3 is a four-step reaction with proton abstraction as the rate-limiting step. Energy decomposition analysis of the transition states reveals that steric hindrance in the phosphine molecule plays a key role in the site-selective formation of the phosphonium salt. The mechanism of 2-Ph-pyridine phosphination with PAr2CF3 is similar to that with PPh3, and analyses of the effects of substituents show that electron-withdrawing groups decreased the nucleophilicity of the phosphine, whereas aryl electron-donating groups increased it. Finally, TfO− plays an important role in the C–H fluoroalkylation of pyridine, as it brings weak interactions.

Keywords: phosphination; metal free; C–H fluoroalkylation; mechanism; DFT

1. Introduction

Organophosphorous compounds have broad applications in the fields of organic synthesis, ligand chemistry, medicinal chemistry, agrochemistry, and materials [1–3]. In light of their fundamental importance, many methods of C–P bond formation have been developed [4–6]. Among these methods, transition-metal-catalyzed cross-coupling reactions, including the Pd- [7], Cu- [8], Zn- [9], Ni- [10], Mn- [11], Ag- [12], and Rh-catalyzed [13] phosphination reactions of various aryl partners with phosphine reagents, have received great attention [14]. In particular, researchers have focused on developing a mild, efficient, and environmentally benign method of C–P bond formation [15–17].

In 1979, the Akiba group succeeded in synthesizing dimethyl phosphonates by reacting quinoline or isoquinoline with acyl chlorides then adding trimethyl phosphite (Scheme 1a) [18]. Since then, many improved methods have been developed. For example, Ander et al. reported the synthesis of PO(OR)2- and PR3-substituted N-heteroaromatic rings by reaction of the N-heteroaromatic ring with phosphorous compounds in the presence of Tf2O and amine (Scheme 1b,c) [19–23]. Although the formation of metal-free C–P bonds has been investigated for many decades, little progress had been made in this area until McNally and co-workers finally succeeded in developing a general approach to form C–O, C–S, C–C, C–D/T, and C–N bonds by transforming pyridines to phosphonium salts then reacting them with nucleophiles (Scheme 1d) [24–32]. The groups of Harutyunyan and Jumde were able to produce pyridyl-ether by directly functionalizing the pyridine ring according to the methodology proposed by McNally and co-workers [33]. The metal-free Sandmeyer-type phosphorylation of aryl amines and electrophilic phosphinative cyclization of alkenes were used to synthesize aryl phosphonates [34] and various phosphine derivatives [35]. Finally, Stephan et al. used Frustrated Lewis Pairs to form compounds with C–P bonds [36].
Despite their importance for the construction of phosphorous compounds and phosphonolates, which can be used as a late-stage functionalization tool, the mechanisms underlying the formation of metal-free C–P bonds are not well understood. McNally et al. proposed a possible reaction pathway for the reaction illustrated in Scheme 1c, as shown in Scheme 1e [27]. According to this mechanism, the reaction is initiated by the nucleophilic attack of pyridine on Tf₂O (I) to form a pyridinium triflyl salt. The addition of phosphine to this salt results in the formation of a C–P bond (II). In the last step, the NEt₃ base abstracts an H-atom from the adduct to restore aromaticity (III). Although this mechanism explains how the C–P bond is formed, it does not account for the factor controlling the site-selective formation of the phosphonium salt. In reaction Scheme 1b, NaI is involved in the generation of the C–P bond, and the product is neutral. Meanwhile, in reaction Scheme 1c, the product is an ion pair. The role of NaI and the reason behind the difference in product nature remain unclear. In this study, we investigate the detailed mechanisms of pyridine reaction with P(OEt)₃, PPh₃, and PAr₂CF₂CXF₂ phosphines, and we assess the effect of the phosphine on the formation of metal-free C–P bonds.

2. Results and Discussion

In this section, we discuss the possible mechanisms of pyridine phosphination with P(OEt)₃, PPh₃, and PAr₂CF₂CXF₂. The site-selectivity of the reaction is also analyzed.

2.1. Mechanism of Pyridine Phosphination with P(OEt)₃

The computed Gibbs free energy profiles corresponding to the phosphination of pyridine with P(OEt)₃ in acetonitrile solvent (Equation (1)) are presented in Figures 1 and 2. The optimized geometries of all stationary points along the reaction pathway are displayed in Figures S1 and S2 of the Supplementary Materials.
1.2 kcal/mol in acetonitrile (relative to pyridine + Tf₂O). The S–O(OTf) bond in TS₁/₂ is elongated to 1.73 Å, whereas the N–S bond distance is decreased to 2.56 Å. The natural population analysis (NPA) charge of the C4 atom of the transition state [37]. The free energy barrier of this reaction step is only 1.73 kcal/mol in acetonitrile (relative to pyridine + Tf₂O), which agrees well with the expected barrier value, considering the experimental conditions (−78 °C to room temperature). The S–O(OTf) bond in TS₁/₂ is elongated to 1.73 Å, whereas the N–S bond distance is decreased to 2.56 Å. The natural population analysis (NPA) charge of the C4 atom of the pyridine moiety increases from −0.159 in free pyridine to −0.06 in intermediate 2, which indicates that this atom becomes more electrophilic along the reaction pathway. Therefore, Tf₂O facilitates the nucleophilic addition reaction by strengthening the electrophilicity of pyridine.

The second reaction step involves the addition of the phosphine P(OEt)₃ nucleophile to intermediate 2 via the TS₃/₄ transition state, yielding N-trifluoromethylsulfonyldihydropyridine.

Figure 1. Free energy profile of pyridine activation and nucleophilic addition.

Figure 2. Free energy profile of dihydropyridine rearomatization to afford the final product (path A).

2.1.1. Activation of Pyridine and Nucleophilic Addition

Based on our calculations, the reaction is initiated by the nucleophilic attack of pyridine on Tf₂O (SN₂ reaction) to form N-(trifluoromethylsulfonyl) pyridium triflate 2 and TfO⁻ via the TS₁/₂ transition state [37]. The free energy barrier of this reaction step is only 1.2 kcal/mol in acetonitrile (relative to pyridine + Tf₂O), which agrees well with the expected barrier value, considering the experimental conditions (−78 °C to room temperature). The S–O(OTf) bond in TS₁/₂ is elongated to 1.73 Å, whereas the N–S bond distance is decreased to 2.56 Å. The natural population analysis (NPA) charge of the C4 atom of the pyridine moiety increases from −0.159 in free pyridine to −0.06 in intermediate 2, which indicates that this atom becomes more electrophilic along the reaction pathway. Therefore, Tf₂O facilitates the nucleophilic addition reaction by strengthening the electrophilicity of pyridine.

The second reaction step involves the addition of the phosphine P(OEt)₃ nucleophile to intermediate 2 via the TS₃/₄ transition state, yielding N-trifluoromethylsulfonyldihydropyridine.
4. This step has a free energy barrier of 11.1 kcal/mol in acetonitrile (relative to 3), which is relatively low, and the C4–P bond distance in TS3/4 is 2.37 Å. The approach of P(OEt)3 distorts the pyridine ring, and in the dearomatized intermediate 4, this ring has a typical dihydropyridine structure.

2.1.2. Rearomatization of Dihydropyridine

The rearomatization of dihydropyridine (intermediate 4) occurs via two possible reaction pathways. The first pathway involves the NaI-mediated migration of an ethyl group, followed by proton abstraction by NEt3 (path A in Figure 2). Meanwhile, in the second pathway, abstraction occurs first, then migration (path B in Figure 3).

![Figure 3. Free energy profile of dihydropyridine rearomatization to afford the final product (path B).](image)

As shown in Figure 2, path A of dihydropyridine rearomatization is mediated by NaI, which can directly extract the ethyl group in 4 via TS5a. However, the free energy barrier of this step is high (40.3 kcal/mol). Alternatively, the TfO⁻ ion in 4 may be exchanged with the I⁻ ion of NaI, which promotes the migration of the ethyl group. The iodide ion may attack the ethyl groups from the frontside via TS6a or from the backside via TS7/8, with free energy barriers of 35.9 and 20.0 kcal/mol, respectively. The attack of I⁻ on the ethyl groups is an S₉2 type reaction, where the iodide ion acts as a nucleophile. Considering that the attack of I⁻ from the backside yields higher orbital overlap than the attack from the frontside (Figure 4), TS7/8 is more favorable than TS5a or TS6a.

![Figure 4. Orbital interactions implicated in the three S₉2 transition states: TS5a, TS6a, and TS7/8.](image)
Subsequently, the product of the $S_N2$ reaction, 4-Phosphonato substituted pyridine 9, reacts with NEt$_3$ to generate intermediate 11 via a proton transfer transition state ($TS_{10/11}$). With a free energy barrier of 13.0 kcal/mol only (relative to 9), this reaction step occurs readily. Intermediate 11 undergoes isomerization to a more stable isomer 12, in which the ammonium cation is close to the P=O group. Then, the N-S bond in 12 breaks to form intermediate 13 via $TS_{12/13}$, which lies at 6.0 kcal/mol above 9. The final product 15 is obtained upon the dissociation of the Et$_3$HN$^+$···CF$_3$SO$_2^-$ ion pair 14 from 13.

In path B, proton abstraction precedes the migration of the ethyl group. The free energy profile of this pathway is shown in Figure 3, and the optimized geometries of all implicated species are given in Figure S3. First, the NEt$_3$ amine abstracts a proton from the C4 of the dihdropyridine intermediate 4 via $TS_{16/17}$ to generate intermediate 17. The free energy barrier of this process is 8.9 kcal/mol (relative to 4), which is relatively low. Then, intermediate 17 isomerizes to a more stable isomer 18, in which the ammonium cation is close to the Tf group. Subsequently, the N-S bond in 18 breaks, resulting in the formation of intermediate 19 via $TS_{18/19}$, which lies at 4.1 kcal/mol above 18. Finally, the iodide ion of NaI abstracts the ethyl group of 19 via an $S_N2$ reaction, whose free energy barrier is only 17.8 kcal/mol ($TS_{21/22}$). This indicates that the I$^-$-mediated migration of the ethyl group is a facile process. In both paths, A and B, ethyl migration is the rate-determining step. The free energy barrier of this step in path B is 17.8 kcal/mol ($TS_{21/22}$), which is comparable to that in path A ($TS_{78}$, 20.0 kcal/mol). This reveals that the reaction can also occur when the order of the addition of NaI and NEt$_3$ in the experiment is reversed.

Overall, our calculations suggest that path A (i.e., the activation of pyridine and nucleophilic addition of P(OEt)$_3$, followed by the migration of the ethyl group and rearomatization of dihdropyridine) is energetically feasible, with an overall energy change of −83.0 kcal/mol. The rate-limiting step is the nucleophilic addition of P(OEt)$_3$ via $TS_{78}$, which has a free energy barrier of 20.0 kcal/mol. Although the order of the elementary steps is reversed in path B, this path can also occur under experimental conditions.

2.1.3. Origin of Site-Selectivity

Based on the experiments conducted in a previous study, the reaction of pyridine with P(OEt)$_3$ results in the nearly exclusive formation of C4-phosphonates (C4-phosphonate:C2-phosphonate = 95:5) [22]. The implicated mechanism of the formation of C2-phosphonate involves nucleophilic addition ($o$-$TS_{3/4}$, 13.7 kcal/mol) and the migration of the ethyl group ($o$-$TS_{7/8}$, 21.9 kcal/mol), as illustrated in Figure 5. The rate-determining step is ethyl group abstraction by I$^-$ ($o$-$TS_{7/8}$). Notably, C2-phosphonates may be formed via the same mechanism as C4-phosphonates; however, the free energy barriers of the key reaction steps are lower in the case of C4-phosphonate formation, as shown in Table 1. Moreover, the energy of the C4-phosphonate product is lower than that of the C2 counterpart. These results indicate that compared to C2-phosphonate formation, the production of C4-phosphonates is more favored both dynamically ($TS_{3/4}$ and $TS_{7/8}$) and thermodynamically (9).

![Figure 5. Free energy profile of pyridine phosphination to afford C2-phosphonates.](image-url)
Table 1. The free energy barriers of key elementary reactions and energies of the C2- and C4-phosphonate products (unit kcal/mol).

| Reaction                        | TS3/4 | TS7/8 | Product (9) |
|---------------------------------|-------|-------|-------------|
| Formation of C4-phosphonate     | 11.1  | 20.0  | −27.2       |
| Formation of C2-phosphonate     | 13.7  | 21.9  | −25.9       |

The C2-phosphonate o-9 undergoes proton abstraction by NEt3 via the o-TS10/11 transition state. The related free energy barrier is 21.9 kcal/mol, compared with 13.1 kcal/mol for proton abstraction from the C4-phosphonate (TS10/11). Considering the relatively high barrier of o-9 rearomatization, this is the main product of the reaction.

To determine the origin of product selectivity, activation strain model (ASM) [38–40] analyses of the TS7/8 and o-TS7/8 transition states were conducted. These transition states may be divided into two parts, one involving phosphorane and I−, and the other involving pyridine and the Tf group. The energy of the interaction between the two parts in TS7/8 and o-TS7/8 is −106.9 and −113.5 kcal/mol, respectively. The larger interaction energy of o-TS7/8 compared to TS7/8 is attributed to the presence of a strong π bond in the former. As for the strain energies of the phosphorane and pyridine moieties, they are larger in o-TS7/8 than in TS7/8 (Table 2), which indicates that the steric hindrance in the former is greater than that in the latter. Considering that the interaction energy and strain energy differences between TS7/8 and o-TS7/8 are −6.6 and 10.8 kcal/mol, respectively, the free energy of o-TS7/8 is larger than that of TS7/8, and the para-substituted phosphonate is the main product. In summary, the product selectivity is determined by the steric hindrance. The phosphination of pyridine with P(OEt)3 to afford C2- and C4-phosphonates was also explored, and the results confirm that steric hindrance plays a key role in site-selectivity. The related free energy profiles and optimized structures are presented in Figures S5–S8.

Table 2. Interaction and strain energies of the TS7/8 and o-TS7/8 transition states (unit kcal/mol).

| TS7/8 | Einter | Estrain |
|-------|--------|---------|
|       | Phosphonate | Pyridine | Sum |
| TS7/8 | −106.9 | 59.8 | 46.1 | 105.9 |
| o-TS7/8 | −113.5 | 67.9 | 48.8 | 116.7 |
| differ | −6.6 | 8.1 | 2.7 | 10.8 |

2.2. Phosphination of Pyridine with PPh3

2.2.1. Mechanism of Pyridine Phosphination with PPh3

As there is no NaI additive, the process of 2-Ph-pyridine phosphination with PPh3 is simpler than that with P(OEt)3 (Equation (2)). The free energy profile of this process is shown in Figure 6, and the geometries of all implicated species are illustrated in Figure S9.

The first and second steps of phosphination with PPh3 are similar to those of phosphination with P(OEt)3, and their free energy barriers are 7.7 (TS23/24) and 7.8 kcal/mol (TS25/26), respectively. Following these steps, the NEt3 amine abstracts a proton from C4 of the dihydropyridine intermediate 26 via the TS27/28 transition state to generate intermediate 28. The free energy barrier of this step is 20.9 kcal/mol (relative to 26). Subsequently, 28 isomerizes produce intermediate 29 and the S–N bond in this intermediate is broken to give the 4-phosphonato substituted pyridine 31 and an ammonium salt 14. Considering that the aromaticity of pyridine is restored by breaking the S–N bond in 29, the free energy barrier of this step is very small (0.4 kcal/mol). Therefore, proton abstraction (TS27/28, 20.9 kcal/mol) is the rate-limiting step in the mechanism of 2-Ph-pyridine phosphination.
with PPh$_3$. The same step in the mechanism of pyridine phosphination with P(OEt)$_3$ has a smaller barrier (TS$_{10/11}$, 13.0 kcal/mol) due to the relatively low steric hindrance imposed by P(OEt)$_3$, whose volume is smaller than PPh$_3$.

Figure 6. Free energy profile of pyridine phosphination with PPh$_3$.

2.2.2. Origin of Site-Selectivity

The experiments conducted in previous studies show that the reaction of 2-Ph-pyridine phosphination with PPh$_3$ is highly selective and favors the formation of the para-substituted product [19,22]. To investigate the reason behind the site-selectivity of this reaction, calculations of 2-Ph-pyridine phosphination at the ortho position were performed. The obtained free energy profile is shown in Figure 7, and the geometries of all implicated species are shown in Figure S10. The nucleophilic attack of PPh$_3$ on the pyridinium salt 24 at the ortho position (TS$_{24/32}$) has a free energy barrier of 7.0 kcal/mol, and the barrier of the subsequent proton abstraction by NEt$_3$ (TS$_{32/33}$) is 22.7 kcal/mol. The latter is higher than the energy barrier of the analogous reaction, leading to the formation of the para-substituted product (TS$_{27/28}$, 20.9 kcal/mol). Based on ASM [38–40] analysis, the difference in the strain energies of the phosphonate moieties in TS$_{27/28}$ and TS$_{32/33}$ are bigger than that of the pyridine and NEt$_3$ moieties (Table 3). This indicates that the difference between the energies of the two transition states is mainly attributed to the phosphonate moiety. Specifically, the steric hindrance at the ortho position of pyridine renders this moiety more distorted. Therefore, the phosphination of pyridine at ortho position is less favorable than that at para position, which agrees well with the available experimental data.

Figure 7. Free energy profile of 2-Ph-pyridine phosphination with PPh$_3$ to give the ortho-substituted product.
Overall, the results indicate that the phosphination of pyridine with PPh₃ may proceed via four successive elementary steps including activation of pyridine, nucleophilic addition, proton abstraction, and S–N bond breaking. The large volume of PPh₃ raises the free energy barrier of proton abstraction. Nevertheless, the reaction is favored by its high exothermicity (−72.7 kcal/mol). The site-selectivity is thus attributed to the unfavorable steric hindrance at the ortho position of pyridine.

2.3. Phosphination of 2-Ph-Pyridine with Diarylfluoroalkylphosphines (PAr₂CF₂X)

Paton and Mcnally previously reported the phosphination of 2-Ph-pyridine with diarylfluoroalkylphosphines (Scheme 2) [41]. They compared five different phosphines and found that the yield of the product correlates with the donating capacity of the phosphine’s aryl substituents.

\[
\text{Ph} + \text{diarylfluoroalkylphosphines} \rightarrow \text{Ph}^{+} + \text{Ar}_{2}R^{2}\text{F}
\]

Scheme 2. Phosphination of 2-Ph-pyridine with diarylfluoroalkylphosphines P1–P5 using DBU as a Lewis base.

Herein, the mechanism of 2-Ph-pyridine phosphination with P1–P5 was studied and the obtained results demonstrate that unlike the phosphination reaction with PPh₃, the rate-determining step is the nucleophilic addition of phosphines. Compared to PPh₃ (proton affinity = 159.4 kcal/mol), the proton affinities of P1–P5 are smaller, as shown in Table 4. This suggests that the CF₃ and CF₂H electron-withdrawing substituents decrease the nucleophilicity of the diarylfluoroalkylphosphines, thereby increasing the energy barrier of nucleophilic addition. As a result, this reaction step becomes the rate-determining step.

Table 3. Strain energies of pyridine, phosphonate and NEt₃ moieties in the TS₂⁷/₂₈ and TS₃₂/₃₃ transition states.

|       | Phosphonate | Pyridine | NEt₃ |
|-------|-------------|----------|------|
| TS₂⁷/₂₈ | 5.9         | 91       | 4.1  |
| TS₃₂/₃₃ | 9.1         | 89.6     | 3.2  |
| differ | 3.2         | −1.4     | −0.9 |

Table 4. Proton affinities of P1–P5 and free energy barriers of the nucleophilic addition reactions of these phosphines to 2-Ph-pyridine.

| Phosphine | R¹       | R²       | Product Yield (%) | ΔGSTS (kcal/mol) | Protonaffinity (kcal/mol) |
|-----------|----------|----------|-------------------|-----------------|-------------------------|
| P1        | H        | CF₃      | n.d.              | 20.1            | 143.9                   |
| P2        | OMe      | CF₃      | 54                | 18.2            | 146.6                   |
| P3        | NMe₂     | CF₃      | 81                | 12.7            | 153.2                   |
| P4        | N-pyrrolidinyl | CF₃ | 85                | 11.1            | 154.3                   |
| P5        | OMe      | CF₂H     | 90                | 11.3            | 153.1                   |

Unlike the electron-withdrawing substituents (CF₃ or CF₂H) of P1–P5, the aryl electron-donating substituents (OMe, NMe₂, and N-pyrrolidinyl) promote the nucleophilicity of the phosphines, as per the proton affinity values listed in Table 4. This means that they...
decrease the barrier of the nucleophilic addition elementary step. Consequently, when the aryl substituent in the diarylfluoroalkylphosphate is an electron-donating group, the nucleophilic addition of this phosphate to 2-Ph-pyridine is facile, and the product yield of the reaction is high. The opposing effects of electron-withdrawing and electron-donating substituents on the rate of 2-Ph-pyridine phosphination with diarylfluoroalkylphosphines agree well with the experimentally observed reactivities of different phosphines [41].

2.4. C-H Fluoroalkylation Reaction of 2-Ph-Pyridine

As mentioned in the introduction, the phosphonium ion can be used as a functional handle to form other chemical bonds. For example, Paton and Mcnally used phosphonium salts to form fluoroalkyl pyridine, and they studied this reaction both, experimentally and theoretically (DFT calculations) [41]. Based on the obtained results, the authors proposed that the reaction process involves water addition and ligand coupling, as shown in Scheme 3, and that the key transition state is the coupling of CF3-PyH+ (CF3-PyH+·TS). In the presence of TfO−, the free energy barrier of this transition state is reduced from 25.8 to 16.2 kcal/mol, which indicates that this anion plays an important role in the CF3-PyH+ coupling reaction. Herein, the interactions in the CF3-PyH+·TfO-TS transition state were analyzed, and as shown in Figure 8, there are two types of weak interactions: hydrogen bonding between TfO− and OH, and dispersion effects between TfO− and the pyridinium ring. These interactions promote ligand coupling.

![Scheme 3](image)

Scheme 3. The fluoroalkylation of pyridine in acidic solvent starting from a phosphonium salt. CF3-PyH+·TS is reported by Paton and Mcnally and CF3-PyH+·TfO-TS is calculated by our group.

![Figure 8](image)

Figure 8. The weak interactions in the CF3-PyH+·TfO-TS transition state.

3. Computational Methods

The Gaussian16 software package [42] was used to perform all DFT calculations according to the self-consistent reaction field (SCRF) method and the IEFPCM solvation model [43], with acetonitrile as solvent. The geometries of all minima and transition states

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were optimized and the harmonic frequencies calculated at the M06-2X/6-311G** level of theory in solution [44–46]. Meanwhile, single-point energy calculations of the minimum energy conformers (i.e., optimum geometries) were performed at the M06-2X/6-311++G** level of theory in solution. The 3D-optimized structures were visualized using the CYLview program [47], and the calculated frequency values were used to correct the free energy at 298.15 K and 1 atm. Based on the “the theory of free volume” [48–53], a correction factor of −2.55 (or 2.55) kcal mol−1 was added to the free energy values to account for the effect of the ideal gas phase model in overestimating the contribution of entropy. Meanwhile, the Gibbs energies were corrected to standard state 1 M [54]. The proton affinity were defined as the negative of the enthalpy change for the reaction (P + H+ → PH+). The important transition states were confirmed using intrinsic reaction coordinate (IRC) analysis [55,56], and the partial atomic charges were allocated based on natural bond orbital (NBO) analyses, which were performed at the M06-2X/6-311G** level [57–59]. To study the electronic structure changes induced by the SN2 reactions along the reaction pathway, intrinsic bond orbital (IBO) analyses were performed [60]. The weak interactions in the transition state were analyzed using the Multiwfn program [61,62].

4. Conclusions

This study uses DFT calculations to elucidate the detailed mechanism of pyridine phosphination with three different phosphines: P(OEt)3, PPh3, and PAr2CF3. As shown in Scheme 4, the reactions are initiated by pyridine activation and nucleophilic addition of phosphine. In the case of phosphination with P(OEt)3, the subsequent steps are NMe2-mediated ethyl migration and rearomatization of dihydropyridine. Meanwhile, in the case of phosphination with PPh3 or PAr2CF3, the rearomatization of dihydropyridine by NEt3 occurs first, followed by ligand coupling of the 4-phosphonato substituted pyridine intermediate to give trifluoromethylated pyridine. Considering that the proton affinity of PAr2CF3 is smaller than the affinities of P(OEt)3 and PPh3, the rate-determining step of pyridine phosphination with PAr2CF3 is nucleophilic addition, whereas that of phosphination with P(OEt)3 or PPh3 is ethyl migration and proton transfer.

Scheme 4. Summary of the possible pathways of pyridine phosphination with P(OEt)3, PPh3, and PAr2CF3.

The steric hindrance of phosphine determines the site-selectivity of the phosphination reaction with P(OEt)3 or PPh3. In the case of PAr2CF3, the effect of the aryl electron-donating substituents increasing the proton affinity and reducing the reaction barrier is greater than the opposing effect of the electron-withdrawing substituents, which facilitates the nucleophilic addition of the phosphine. These electronic and steric effects on the rate of the reaction further support our proposed mechanism. Finally, TIO− plays an important role in the C–H fluoroalkylation of 2-Ph-pyridine, as it induces hydrogen bonding interactions and dispersion effects in the ligand coupling transition state.
Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/molecules27175694/s1, Figure S1: Optimized intermediates and transition states of pyridine activation and nucleophilic addition; Figure S2: Optimized intermediates and transition states of dihydropyridine re aromatization to afford the final product (path A); Figure S3: Optimized intermediates and transition states of dihydropyridine re aromatization to afford the final product (path B); Figure S4: Optimized intermediates and transition states of pyridine phosphination to afford C2-phosphonates; Figure S5: Optimized intermediates and transition states of phosphination of pyridine with \( \text{P(OiPr)}_3 \) to afford C4-phosphonates; Figure S6: Optimized intermediates and transition states of phosphination of pyridine with \( \text{P(OiPr)}_3 \) to afford C4-phosphonates; Figure S7: Free energy profile of phosphination of pyridine with \( \text{P(OiPr)}_3 \) to afford C4-phosphonates; Figure S8: Free energy profile of phosphination of pyridine with \( \text{P(OiPr)}_3 \) to afford C2-phosphonates; Table S1: Interaction and strain energies of the ipr-p-TS\(_{7/8}\) and ipr-o-TS\(_{7/8}\) transition states (unit kcal/mol); Figure S9: Optimized intermediates and transition states of pyridine phosphination with \( \text{PPh}_3 \); Figure S10: Optimized intermediates and transition states of 2-Ph-pyridine phosphination with \( \text{PPh}_3 \) to give the ortho-substituted product; Figure S11: Optimized intermediates and transition states of Phosphination of 2-Ph-pyridine with diarylfluoroalkylphosphines P1–P5; Figure S12: Optimized intermediates and transition states of The fluoroalkylation of pyridine in acidic solvent starting from a phosphonium salt; Table S2: Corrected free energies of all species; Table S3: Imaginary frequencies of all transition states; Table S4: Cartesian coordinates of all species.

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