Electrophilic Activation of Amides for the Preparation of Poly-substituted Pyrimidines

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Abstract In this article we describe the straightforward synthesis of polysubstituted pyrimidines by electrophilic activation of secondary amides in the presence of alkynes. An unusual mechanistic detour leading to pyridine derivatives as products is also presented and briefly discussed.

Key words pyrimidines, amide activation, hydride shift, pyridines

Heteroaromatics occupy a prominent role in organic chemistry. Among them, the pyrimidine core is present in several natural products, pharmaceuticals and functional materials. For these reasons, numerous methods have been developed throughout the years to prepare these motifs (Scheme 1). The most popular method for the synthesis of pyrimidines remains the condensation of amidines with carbonyls (Scheme 1, a).4,5 Movassaghi and co-workers developed a straightforward synthesis of pyrimidines by electrophilic activation of a secondary enamide with triflic anhydride (Scheme 1, c).6 In that elegant report, the requirement for an electron-rich alkene in the form of the enamide partner somewhat limits the scope of the reaction. We herein propose an approach consisting of a three-component reaction that merges one alkyne and two secondary amides under an electrophilic activation regime (Scheme 1, d).

We started our investigation of this transformation by employing N-cyclopentylbenzamide (1a), 2-iodopyridine as a base and prop-1-yn-1-ylbenzene as alkyne partner (Table 1). Already on our first trial (Table 1, entry 1) using 3 equivalents of amide, base and triflic anhydride, the expected pyrimidine 2a was obtained in 60% NMR yield. Other halogenated bases were screened, such as 2-fluoropyridine and 2-chloropyridine (Table 1, entries 2 and 3), resulting in an increase to 77% and 81% yield, respectively. During these experiments, we became aware of the non-negligible effect of the quenching method employed. In the event, we found that stirring for 1 hour at room temperature with a saturated aqueous solution of sodium bicarbonate increased the yield (Table 1, entry 4), up to 88% with 2-iodopyridine. Further variation of the base (Table 1, entry 5) or the loading of activated amide (Table 1, entry 6) led only to lower yields. Finally, we decided to modify the N-amide substituent and investigated N-isopropyl- and N-propylbenzamide. Both were less effective than 1a (yielding 57% and 0%, respectively; cf. Table 1, entries 7 and 8).

Another approach which has gained traction in recent years is the metal-catalyzed [2+2+2] cycloaddition between an alkyne and two nitriles (Scheme 1, b).4 Recently, our group reported a metal-free version of this reaction using ynamides and relying on triflic acid for the activation of the C≡C bond.7 In that elegant report, the requirement for an electron-rich alkene in the form of the ynamide partner somewhat limits the scope of the reaction. We herein propose an approach consisting of a three-component reaction that merges one alkyne and two secondary amides under an electrophilic activation regime (Scheme 1, d).

We started our investigation of this transformation by employing N-cyclopentylbenzamide (1a), 2-iodopyridine as a base and prop-1-yn-1-ylbenzene as alkyne partner (Table 1). Already on our first trial (Table 1, entry 1) using 3 equivalents of amide, base and triflic anhydride, the expected pyrimidine 2a was obtained in 60% NMR yield. Other halogenated bases were screened, such as 2-fluoropyridine and 2-chloropyridine (Table 1, entries 2 and 3), resulting in an increase to 77% and 81% yield, respectively. During these experiments, we became aware of the non-negligible effect of the quenching method employed. In the event, we found that stirring for 1 hour at room temperature with a saturated aqueous solution of sodium bicarbonate increased the yield (Table 1, entry 4), up to 88% with 2-iodopyridine. Further variation of the base (Table 1, entry 5) or the loading of activated amide (Table 1, entry 6) led only to lower yields. Finally, we decided to modify the N-amide substituent and investigated N-isopropyl- and N-propylbenzamide. Both were less effective than 1a (yielding 57% and 0%, respectively; cf. Table 1, entries 7 and 8).
Biographical Sketches

Dr. Tobias Stopka was born in Münster, Germany in 1988. In 2011 he obtained his Bachelor’s degree and in 2013 his Master’s degree in organic chemistry from the Westfälische Wilhelms University Münster. After having moved to Aachen, he started his doctoral studies in the group of Prof. Dr. Meike Niggemann at the RWTH Aachen (Germany) working on reactions based on vinyl cations as reactive intermediates. In 2017, he successfully defended his PhD and graduated with honors. For his dissertation he was awarded the Borchers Badge. Until 2018 he was a postdoctoral researcher in the group of Prof. Nuno Maulide in Vienna (Austria) where he worked on sulfur ylide based transformations and amide activation. Currently, he is working as a business consultant.

Dr. Pauline Adler was born in 1988 in Metz, France. In 2010 she obtained her Bachelor’s degree in physical and molecular chemistry from the Université Paris-Sud. In 2012 she passed the ‘Agrégation’ of Physics and Chemistry with success. She obtained her Master’s degree from the Université Paris-Sud in 2013 with honors. Thanks to a ‘Allocation Spécifique pour Normalien’ doctoral research grant from the French Ministry of Higher Education and Research, she started a PhD in the group of Dr. Nicolas Rabasso on the study of the synthesis and reactivity of unsaturated aminophosphonates. She defended her PhD in 2016. Since 2017, she has been working as a postdoctoral researcher in the group of Prof. Maulide in Vienna (Austria). Her work is focused on the chemoselective activation of amides.

Gerhard Hagn was born in 1991 in Schärding, Austria. He obtained his Bachelor’s degree in chemistry working on molecular dynamics simulations of ionic liquids at the University of Vienna (Austria), Department of Computational Biological Chemistry. To complete his M.Sc. degree, he joined the group of Prof. Christopher Gerner at the University of Vienna, Department of Analytical Chemistry.

Haoqi Zhang was born in Baden, Austria in 1994. He obtained his Bachelor’s degree in chemistry from the University of Vienna (Austria) in 2017 by working on synthetic studies towards the total synthesis of virol A, in the group of Prof. Maulide. His interests lie in synthetic chemistry and he will continue his Master’s studies in the field of organic and biological chemistry at the University of Vienna.

Veronica Tona was born in 1990 in Loreto, Italy. She obtained her Bachelor’s and Master’s degrees from the University of Camerino (Italy) under the supervision of Prof. Marcantoni. Since 2015, she has been a PhD student in the group of Prof. Maulide in Vienna (Austria). Her research interests include electrophilic amide activation and metal-free amination reactions.

Prof. Nuno Maulide was born in 1979 in Lisbon, Portugal and since 2013 has been a Full Professor and Chair of Organic Synthesis at the University of Vienna (Austria). Prior to that, he was a Max Planck Group Leader at the Max-Planck-Institut für Kohlenforschung (Mülheim, Germany) from 2009, following PhD studies at the Université catholique de Louvain, Belgium (Prof. I. E. Markó) and a postdoctoral stay at Stanford University, USA (Prof. B. M. Trost). Prof. Maulide is a regular speaker at international symposia and conferences, having delivered over 20 invited/plenary lectures per year since 2012. His research interests revolve around the exploration of rearrangements and high-energy intermediates in organic chemistry, and have been acknowledged by several awards (including the Bayer Early Excellence Award 2012, the Heinz Maier-Leibnitz Prize 2013, the EurJOC Young Researcher Award 2015, the Elisabeth Lutz Prize of the Austrian Academy of Sciences 2016, the Springer Heterocyclic Chemistry Award 2018, and three ERC Grants – StG 2011, CoG 2016 and PoC 2018). He was elected corresponding member of the Austrian Academy of Sciences in 2018.
Triflic anhydride is well-known to chemoselectively activate tertiary and secondary amides.\textsuperscript{7,8} We propose the following mechanism to explain the observed reactivity (Scheme 2). The secondary amide 1 is activated as nitrilium ion RI-1, reversibly stabilized by 2-iodopyridine (cf. RI-2).\textsuperscript{7b,9} This pivotal nitrilium species can evolve either to the nitrile 3 by loss of cyclopentyl carbocation (presumably favoured by high temperatures), or undergo addition of the alkyne reactant to generate the vinyl cation RI-3. The latter is stabilized by the vicinal aryl group, accounting for the regioselectivity observed in this process. Interception of RI-3 by nitrile 3 (cf. RI-4) sets the stage for cyclization to the pyrimidinium ion RI-5. A second elimination of cyclopentyl carbocation finally accounts for the formation of the pyrimidine product.\textsuperscript{10}

For the investigation of the amide scope, prop-1-yn-1-ylbenzene was retained as the alkyne partner (Scheme 4). The arylamide can be substituted ortho or meta (2\texttextsuperscript{h} and 2\texttextsuperscript{i}), and the presence of halogens on the aromatic ring is also well-tolerated (2\texttextsuperscript{j}), including fluorine (2\texttextsuperscript{k}, 90% yield). A benzamide is not a prerequisite for successful reaction and it was possible to prepare pyrimidine 2\texttextsuperscript{m} starting from a tert-

### Table 1 Optimization of the Reaction\textsuperscript{a}

<table>
<thead>
<tr>
<th>Entry</th>
<th>R</th>
<th>Amide</th>
<th>2-X-Pyr</th>
<th>Tf\textsubscript{2}O</th>
<th>Comment</th>
<th>NMR yield\textsuperscript{b}</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>(\text{1a})</td>
<td>3 equiv</td>
<td>2-I-Pyr (3 equiv)</td>
<td>3 equiv</td>
<td>–</td>
<td>60%</td>
</tr>
<tr>
<td>2</td>
<td>3 equiv</td>
<td>2-F-Pyr (3 equiv)</td>
<td>3 equiv</td>
<td>–</td>
<td>77%</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>3 equiv</td>
<td>2-Cl-Pyr (3 equiv)</td>
<td>3 equiv</td>
<td>–</td>
<td>81%</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>3 equiv</td>
<td>2-I-Pyr (3 equiv)</td>
<td>3 equiv</td>
<td>quench 1 h, NaHCO\textsubscript{3}</td>
<td>88% (83%\textsuperscript{c})</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>3 equiv</td>
<td>2-Cl-Pyr (3 equiv)</td>
<td>3 equiv</td>
<td>quench 1 h, NaHCO\textsubscript{3}</td>
<td>79%</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>2.5 equiv</td>
<td>2-I-Pyr (2.5 equiv)</td>
<td>2.5 equiv</td>
<td>quench 1 h, NaHCO\textsubscript{3}</td>
<td>69%</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>3 equiv</td>
<td>2-I-Pyr (3 equiv)</td>
<td>3 equiv</td>
<td>quench 1 h, NaHCO\textsubscript{3}</td>
<td>57%</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>3 equiv</td>
<td>2-I-Pyr (3 equiv)</td>
<td>3 equiv</td>
<td>quench 1 h, NaHCO\textsubscript{3}</td>
<td>0%</td>
<td></td>
</tr>
</tbody>
</table>

\textsuperscript{a} Reaction conditions: amide, 2-halopyridine, 1,2-dichloroethane (3 mL), 0 °C; addition of Tf\textsubscript{2}O, after 15 min, addition of alkyne (0.2 mmol, 1 equiv); 90 °C, 18 h.

\textsuperscript{b} NMR yield calculated using 1,3,5-trimethoxybenzene as internal standard.

\textsuperscript{c} Isolated yield.

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**Scheme 2** Mechanistic proposal for the pyrimidine synthesis reported herein
butyl amide. Alkenyl amides and benzamides carrying electron-withdrawing substituents led to lower yields, presumably due to less efficient electrophilic activation (2n, 2o).

In the course of this study, we noticed that no pyrimidine was formed when particularly electron-rich alkynes were employed. Instead, as depicted in Scheme 5, a pyridine derivative was isolated. Indeed, from 1-ethynyl-4-methoxybenzene, pyridine 3a was obtained in a moderate 45% yield. A propargylsilane similarly led to a (desilylated) pyridine (3b). Such cycloannulated pyridines can be found in agrochemicals and pharmaceuticals, and are commonly prepared by condensation reactions, in particular the Friedländer annulation.

This unexpected reactivity is rationalized by invoking a 1,5-hydride shift on the intermediate RI-3 to form the carbocation RI-6/azoniaallene RI-6'. Deprotonation thereof yields the imine RI-7 poised for a 6π-electrocyclization step towards the dihydropyridine 4. We presume that 4 undergoes disproportionation to tetrahydropyridine (also detected by HRMS) and pyridine 3, the latter being the only product that can be isolated from the reaction mixture. It is
tempting to presume that the aforementioned (cf. Scheme 2) nitrile capture of RI-3 dominates for ‘less-stabilized’ versions of this intermediate, whereas the intramolecular hydride-transfer pathway is favoured by increased stabilization (and thus increased lifetime in solution) of RI-3.

In conclusion, we have developed a new and efficient access to pyrimidines by formal cycloaddition of 2 equivalents of an appropriately substituted amide and an alkyne. We believe this work is complementary to other methods for the preparation of pyrimidines and benefits from the ready availability of both starting materials. Furthermore, an unusual pathway towards pyridines was uncovered, presumably relying on an internal hydride transfer.

All glassware was flame-dried before use and all reactions were performed under an atmosphere of argon. All solvents were distilled from appropriate drying agents prior to use. Triflic anhydride (Tf₂O) was distilled over P₂O₅ prior to use. All other reagents were used as received from commercial suppliers unless otherwise stated. Reaction progress was monitored by TLC performed on aluminium plates coated with silica gel 60F₂₅₄ with 0.2 mm thickness. Chromatograms were visualized by fluorescence quenching with UV light at 254 nm or by staining using potassium permanganate. Flash column chromatography was performed using silica gel 60 (230–400 mesh, Merck and Co.). Neat IR spectra were recorded using a Perkin-Elmer Spectrum 100 FT-IR spectrometer. Wavenumbers are reported in cm⁻¹.

Mass spectra were obtained using a Finnigan MAT 8200 (70 eV) or an Agilent 5973 (70 eV) spectrometer, using electrospray ionization (ESI). All ¹H NMR, ¹³C NMR and ¹⁹F NMR spectra were recorded using a Bruker AV-400, AV-600 or AV-700 spectrometer at 300 K. Chemical shifts are given in parts per million (ppm, δ), referenced to the solvent peak of CDCl₃, defined at δ = 7.26 ppm (¹H NMR) and δ = 77.16 ppm (¹³C NMR). Coupling constants (J) are quoted in Hz. ¹H NMR splitting patterns are designated as singlet (s), doublet (d), triplet (t), quartet (q), pentet (p). Splitting patterns that could not be interpreted or easily visualized are designated as multiplet (m) or broad (br). Selected ¹³C NMR spectra were recorded using the attached proton test (APT) to facilitate the confirmation and assignment of the structure.

**Synthesis of Amides**

**General Procedure A**

To a solution of the amine (1.00 equiv) and Et₃N (1.50 equiv) in DCM (0.1 M) at 0 °C, the corresponding acyl chloride (1.20 equiv) was added dropwise and the resulting reaction mixture was allowed to warm to room temperature overnight (14 h). After this time, the thionyl chloride and solvent were removed under reduced pressure to afford the acyl chloride. To a solution of the amine (1.00 equiv) and Et₃N (1.50 equiv) in DCM (0.1 M) at 0 °C, the corresponding acyl chloride (1.00 equiv) was added dropwise and the resulting reaction mixture was allowed to warm to room temperature while stirring overnight (14 h). After this time, saturated aqueous NaHCO₃ solution was added and the biphasic system was separated. The aqueous phase was extracted with DCM (1 ×) and the organic phases were combined and dried over anhydrous Na₂SO₄. The dried solution was filtered and concentrated under reduced pressure. The resulting crude material was filtered and concentrated under reduced pressure. The resulting crude material was purified by flash column chromatography on silica gel (heptane/EtOAc) to afford the desired compound.

**Procedure C**

To a solution of the carboxylic acid (1.00 equiv) and DMF (1 drop) in DCM (0.1 M) was added dropwise thionyl chloride (2.00 equiv) at room temperature. The resulting solution was stirred at room temperature overnight (14 h). After this time, the thionyl chloride and solvent were removed under reduced pressure to afford the acyl chloride. To a solution of the amine (1.00 equiv) and Et₃N (1.50 equiv) in DCM (0.1 M) at 0 °C, the corresponding acyl chloride (1.00 equiv) was added dropwise and the resulting reaction mixture was allowed to warm to room temperature while stirring overnight (14 h). After this time, saturated aqueous NaHCO₃ solution was added and the biphasic system was separated. The aqueous phase was extracted with DCM (1 ×) and the organic phases were combined and dried over anhydrous Na₂SO₄. The dried solution was filtered and concentrated under reduced pressure. The resulting crude material was purified by flash column chromatography on silica gel (heptane/EtOAc) to afford the desired compound.

**N-Cyclopentylbenzamide (1a)**

General Procedure A; 97% yield. All analytical data were in good accordance with reported data.¹⁵

**N-Isopropylbenzamide**

General Procedure A; quant. All analytical data were in good accordance with reported data.¹⁶

**N-Propylbenzamide**

General Procedure A; quant. All analytical data were in good accordance with reported data.¹⁷

**N-Cyclopentyl-2-methylbenzamide (1b)**

General Procedure A; quant. All analytical data were in good accordance with reported data.¹⁸

**N-Cyclopentyl-3-methylbenzamide (1i)**

General Procedure A; 95% yield. All analytical data were in good accordance with reported data.¹⁹

**2-Bromo-N-cyclopentylbenzamide (1j)**

General Procedure A; light-yellow solid, quant.; mp 96–98 °C.

IR (neat): 3244, 3066, 2952, 2866, 1628, 1593, 1541, 1468, 1430, 1360, 1320, 1277, 1259, 1187, 1047, 754, 729, 694 cm⁻¹.

¹H NMR (600 MHz, CDCl₃): δ = 7.56 (dd, J = 8.0, 1.0 Hz, 1 H), 7.52 (dd, J = 7.6, 1.7 Hz, 1 H), 7.34 (td, J = 7.5, 1.1 Hz, 1 H), 7.26–7.22 (m, 1 H), 5.93 (br s, 1 H), 4.46–4.37 (m, 1 H), 2.10–2.03 (m, 2 H), 1.75–1.63 (m, 4 H), 1.58–1.52 (m, 2 H).

¹³C NMR (151 MHz, CDCl₃): δ = 167.2, 138.2, 133.4, 131.2, 129.8, 127.7, 119.4, 52.0, 33.1, 23.9.

HRMS (ESI⁺): m/z [M + H⁺] calc for C₁₁H₁₁BrNO: 268.0332; found: 268.0336.
N-Cyclopentyl-4-fluorobenzamide (1k)
[CAS Reg. No. 300829-28-1]
General Procedure A; light-yellow solid, 96% yield; mp 133–135 °C.
IR (neat); 3277, 3077, 2957, 2870, 1719, 1632, 1541, 1435, 1368, 1309, 1304, 1295, 1292, 1285, 1284, 1283.5, 1233, 17.9.
HRMS (ESI+): m/z [M + Na]+ calcd for C_{23}H_{19}N_{2}Na: 379.0972; found: 379.1090.

N-Cyclopentyl-4-methoxybenzamide (1l)
General Procedure A; 90% yield. All analytical data were in good accordance with reported data.20

N-Cyclopentylvalalamide (1m)
General Procedure A; 92% yield. All analytical data were in good accordance with reported data.21

N-Cyclopentylcinnamamide (1n)
General Procedure A; 92% yield. All analytical data were in good accordance with reported data.22

Methyl 4-(Cyclopentylcarbamoyl)benzoate (1o)
[CAS Reg. No. 1325090-17-2]
Procedure C; light-yellow solid, 85% yield; mp 172–174 °C.
IR (neat); 3277, 3077, 2957, 2870, 1719, 1632, 1541, 1435, 1368, 1309, 1304, 1295, 1292, 1285, 1284, 1283.5, 1233, 17.9.
HRMS (ESI+): m/z [M + Na]+ calcd for C_{23}H_{19}N_{2}Na: 379.0972; found: 379.1090.

Synthesis of Pyrimidines 2a–o; General Procedure
All reactions were run on a 0.2 mmol scale.

To a solution of amide (0.6 mmol, 3 equiv) and 2-iodopyridine (2-I-Pyr, 3 equiv) in 1,2-dichloroethane (3 mL), Tf2O (3 equiv) was added at 0 °C. The reaction mixture was stirred for 15 min at 0 °C. The alkyne (1 equiv) was then added and the reaction mixture was stirred at 90 °C for 18 h. After cooling to room temperature, the reaction was quenched with saturated aqueous NaHCO3 solution (3 mL) and the mixture was stirred for 1 h. Then, the aqueous layer was extracted with DCM. The combined organic layers were dried over MgSO4 and the solvent was removed under reduced pressure. The crude residue was purified by flash column chromatography on silica gel (eluents: heptane).

5-Methyl-2,4,6-triphenylpyrimidine (2a)
Yield: 83%.

5-Ethyl-2,4,6-triphenylpyrimidine (2b)
Yield: 72%.

5-(2-Chlorophenyl)-5-methyl-2,6-diphenylpyrimidine (2e)
Yield: 41%.

5-Ethyl-2,4,6,7-tetraphenylpyrimidine (2d)
Yield: 41%.

4-(2-Chlorophenyl)-5-methyl-2,6-diphenylpyrimidine (2f)
Yield: 59%.

4-(4-Chlorophenyl)-5-methyl-2,6-diphenylpyrimidine (2g)
Yield: 58%.

4-(3-Chlorophenyl)-5-methyl-2,6-diphenylpyrimidine (2h)
Yield: 59%.

4-(4-Chlorophenyl)-5-methyl-2,6-diphenylpyrimidine (2i)
Yield: 58%.

4-(4-Chlorophenyl)-5-methyl-2,6-diphenylpyrimidine (2j)
Yield: 59%.

4-(4-Chlorophenyl)-5-methyl-2,6-diphenylpyrimidine (2k)
Yield: 59%.

4-(4-Chlorophenyl)-5-methyl-2,6-diphenylpyrimidine (2l)
Yield: 59%.

4-(4-Chlorophenyl)-5-methyl-2,6-diphenylpyrimidine (2m)
Yield: 59%.

4-(4-Chlorophenyl)-5-methyl-2,6-diphenylpyrimidine (2n)
Yield: 59%.
2,4-Bis(2-bromophenyl)-5-methyl-6-phenylpyrimidine (2j)

Yellow solid, 59% yield; mp 141–143 °C.

IR (neat): 3055, 2926, 2857, 2226, 1557, 1534, 1391, 906, 755, 726, 712 cm\(^{-1}\).

HRMS (ESI\(^+\)): m/z [M + Na\(^+\)]\(^+\) calcd for C\(_{24}\)H\(_{15}\)BrN\(_2\): 478.9753; found: 478.9752.

4-(4-Fluoro-2-methylphenyl)-2,6-diphenylpyrimidine (2g)

IR-light solid; mp 119–120 °C.

IR (neat): 3068, 2930, 2930, 2830, 1502, 1497, 1361, 1248, 1179, 1179, 1072 cm\(^{-1}\).

HRMS (ESI\(^+\)): m/z [M + Na\(^+\)]\(^+\) calcd for C\(_{25}\)H\(_{22}\)N\(_2\)Na: 373.1675; found: 373.1673.

5-Methyl-4-phenyl-2,6-diphenylpyrimidine (2h)

Light yellow solid, 70% yield; mp 182–185 °C.

IR (neat): 3068, 2930, 2930, 2830, 1502, 1497, 1361, 1248, 1179, 1179, 1072 cm\(^{-1}\).

HRMS (ESI\(^+\)): m/z [M + Na\(^+\)]\(^+\) calcd for C\(_{25}\)H\(_{22}\)N\(_2\)NaO\(_2\): 405.1570; found: 405.1573.

2,4-Bis(4-fluorophenyl)-5-methyl-6-phenylpyrimidine (2k)

Yellow solid, 90% yield; mp 148–150 °C.

IR (neat): 3057, 3025, 2928, 2861, 1713, 1601, 1532, 1507, 1392, 1379, 1222, 1149, 1006, 842, 808, 772, 734, 697 cm\(^{-1}\).

HRMS (ESI\(^+\)): m/z [M + Na\(^+\)]\(^+\) calcd for C\(_{25}\)H\(_{22}\)N\(_2\)Br: 478.9753; found: 478.9752.
**Dimethyl 4,4′-(5-Methyl-6-phenylpyrimidine-2,4-diyl)dibenzoate (2o)**

White solid, 21% yield; mp 188–190 °C.

IR (neat): 3057, 2946, 2841, 1595, 1577, 1596, 1561, 1492, 1484, 1440, 1437, 1377, 1371, 1250, 1177, 1132 cm⁻¹.

1H NMR (600 MHz, CDCl₃): δ = 7.38–7.34 (m, 1 H), 7.29 (s, 1 H), 3.09 (t, J = 7.7 Hz, 2 H), 2.90 (t, J = 7.4 Hz, 2 H), 2.31 (s, 3 H), 2.20–2.11 (m, 2 H).

13C NMR (151 MHz, CDCl₃): δ = 165.4, 156.3, 143.4, 140.3, 134.9, 128.7, 128.4, 127.1, 119.6, 34.7, 29.1, 22.7, 19.2.


**Synthesis of Pyridines 3a and 3b**

All reactions were run on a 0.2 mmol scale, using the same general procedure as for the synthesis of pyrimidines 2a-o.

**4-(4-Methoxyphenyl)-2-phenyl-6,7-dihydro-5H-cyclopenta[b]pyridine (3a)**

Light-yellow solid, 63% yield.

IR (neat): 3057, 2946, 2841, 1595, 1577, 1457, 1437, 1424, 1377, 1228, 1076, 1028, 864 cm⁻¹.

1H NMR (600 MHz, CDCl₃): δ = 7.94–7.92 (m, 2 H), 7.46–7.42 (m, 2 H), 7.38–7.34 (m, 1 H), 7.29 (s, 1 H), 3.09 (t, J = 7.7 Hz, 2 H), 2.90 (t, J = 7.4 Hz, 2 H), 2.31 (s, 3 H), 2.20–2.11 (m, 2 H).

13C NMR (151 MHz, CDCl₃): δ = 165.4, 156.3, 143.4, 140.3, 134.9, 128.7, 128.4, 127.1, 119.6, 34.7, 29.1, 22.7, 19.2.


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**Supporting Information**

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**References**


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(10) It is likely that the 2 equivalents of cyclopentyl carbocation eliminated ultimately lead to cyclopentene by proton loss (E1). Given the volatility of cyclopentene, it was not possible to detect its presence in the reaction mixture.


(13) Dihydropyridine 4 has been detected by HRMS in several reactions but was never isolated.


