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**Reference:**

Zhu Yan-Ping, Sergejev Sergey, Franck Philippe, Orru Romano V. A., Maes Bert.- Amine activation : synthesis of N-(hetero)arylamides from isothioureas and carboxylic acids

Organic letters / American Chemical Society - ISSN 1523-7060 - 18:18(2016), p. 4602-4605

Full text (Publisher's DOI): <http://dx.doi.org/doi:10.1021/ACS.ORGLETT.6B02247>

To cite this reference: <http://hdl.handle.net/10067/1348370151162165141>

# Amine activation: Synthesis of *N*-(Hetero)arylamides from Isothioureas and Carboxylic Acids.

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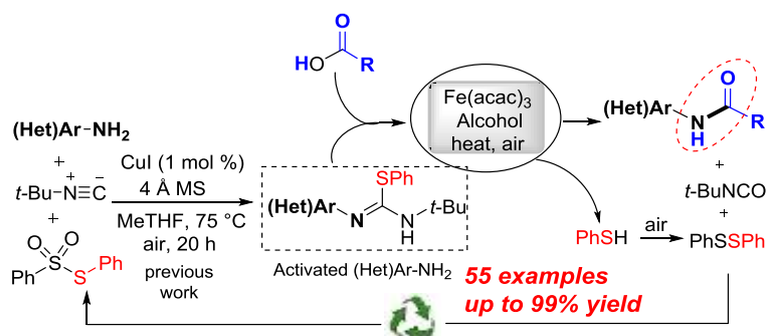
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## Abstract:

A novel method for *N*-(hetero)arylamide synthesis based on rarely explored amine activation, rather than classical acid activation, is reported. The activated amines are easily prepared using a

three component reaction with commercial reagents. The new method shows a broad scope including challenging amides, not (efficiently) accessible via classical protocols.



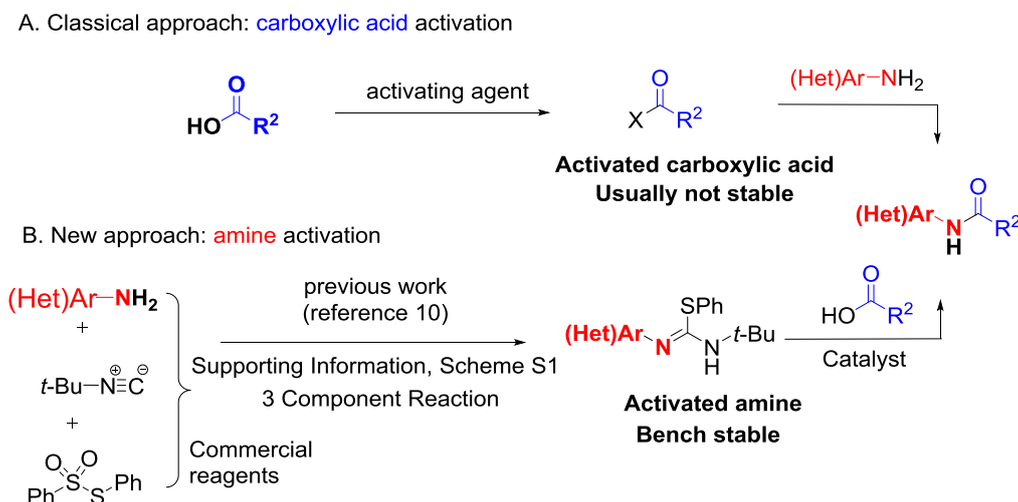
The synthesis of the amide functional group is paramount in organic chemistry. This structural motif is truly ubiquitous, and is present in (bio)polymers (e.g. Nylon, proteins), peptides, natural products (e.g. paclitaxel, penicillin), ligands for catalysis, synthetic drugs and agrochemicals.<sup>1</sup> The main classical route for its synthesis is based on the activation of a carboxylic acid through formation of an acyl chloride, acyl imidazole, anhydride or activated ester, followed by reaction with an amine.<sup>2</sup> The formation of activated esters with phosphonium/uronium/guanidinium salts, based on (azo)benzotriazole derivatives such as HOBt or HATU, is currently the most successful and popular method. However, due to their explosive nature, transportation and storage issues resulted in the development of (1-cyano-2-ethoxy-2-oxoethylideneaminoxy)dimethylamino-morpholino-carbenium hexafluorophosphate (COMU).<sup>3</sup> The prevalence of the amide functionality and the plethora of coupling reagents available for carboxylic acid activation might give the impression that

amide synthesis is a mature transformation. However, this is incorrect as practice proves that often even simple amides resist formation. This triggers the development of always new and more complex and expensive coupling reagents. More recently, Lewis acids based on boron species or metal salts have been reported as suitable catalysts, allowing direct coupling of amine and carboxylic acid when water is simultaneously removed.<sup>4</sup> The difficulty to drive reactions to completion and limited catalyst stability restrict the take up of this technology in industry. Although all these methods cited are very useful for the synthesis of a variety of amides, there are still limitations with respect to amide scope (electronics and sterics), functional group tolerance and chemoselectivity towards other nucleophiles, which require upfront protection thus reducing the efficiency of the process. Based on these limitations many interesting alternative procedures have been developed from precursors other than amines and/or carboxylic acids.<sup>5a-s</sup> However, procedures starting from carboxylic acids and amines are still the most interesting, due to the widespread availability of these building blocks. For challenging amides, derived from sterically hindered carboxylic acids and electron deficient and/or sterically hindered (hetero)aromatic amines, for instance, the classical approaches give a low yield or often even fail, and no general efficient synthetic method is currently at hand. This is illustrated by the synthesis of *N*-mesityl-2,4,6-trimethylbenzamide from 2,4,6-trimethylbenzoic acid and 2,4,6-trimethylaniline (Supporting Information, SI, Scheme S2). A variety of classical coupling reagents under different conditions were tested but generally no reaction product was formed, only with PyBOP a moderate yield was obtained.<sup>6a-f</sup> These poor results and the frequent occurrence of the *N*-(hetero)arylamide moiety in active ingredients (AI) stimulated us to develop a new synthetic method. The importance of this amide subclass can be illustrated by the list of FDA-approved drugs of 2015 and the top 100 drugs based on prescription. In the former list 15% and in the latter 9% of the drugs feature the *N*-(hetero)arylamide entity.<sup>7a-b</sup> Top selling agrochemicals, such as the fungicides Boscalid and Fluxapyroxad, often also contain such a structural moiety.<sup>8</sup>

We initially focused our efforts on the development of a new synthetic protocol for *N*-(hetero)arylamide synthesis, as an example of challenging amide synthesis. Our new approach involves (hetero)arylamine activation by transformation into the corresponding isothiourea, followed by reaction with carboxylic acid under base metal catalysis (Scheme 1).<sup>9</sup> The activation of (hetero)arylamine can be done in high yield in one step by using our recently developed three component reaction (SI, Scheme S1).<sup>10</sup> The reagents (*t*-butyl isocyanide and *S*-phenyl benzenethiosulfonate) required for amine activation are commercially available. While classical synthetic approaches for amide synthesis involve carboxylic acid activation (C→N direction) few procedures are based on amine activation (N

→ C direction: isocyanide,<sup>11</sup> imidazolylcarbonyl,<sup>12</sup> iminophosphorane<sup>13</sup>) (Scheme 1). Moreover, isothiurea are bench stable compounds while activated carboxylic acids are usually unstable and have to be generated *in-situ*.

**Scheme 1:** Classical (A) and new strategy (B) for *N*-(hetero)arylamide synthesis from (hetero)aromatic amines and carboxylic acids

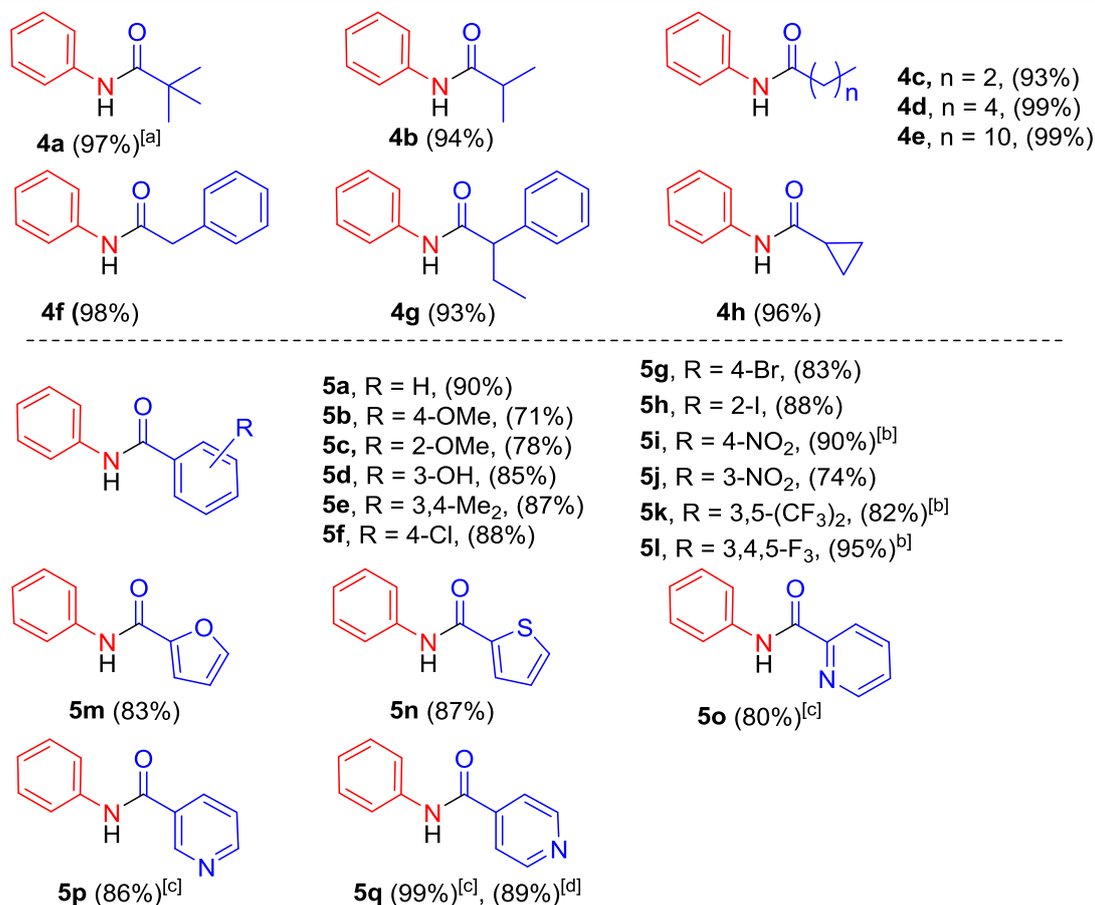
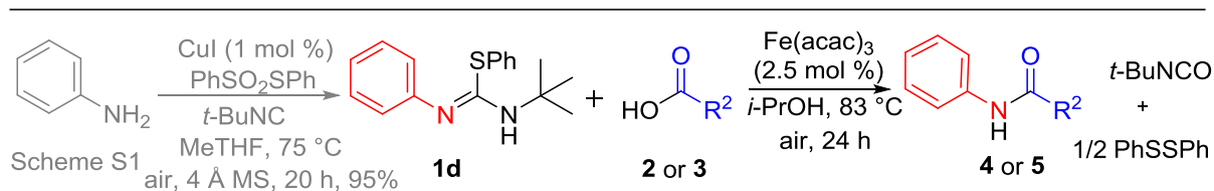


Optimization of the reaction conditions was done using *N*-*t*-butyl-*S*-methyl-*N*-phenylisothiurea (**1a**), as activated aniline, and pivalic acid (**2a**) as model reagents. A catalyst screening in DMF revealed that Fe(acac)<sub>3</sub> was optimal (SI, Table S1). Further screening showed that the reaction worked well in isopropanol (93% yield) when 1.2 equivalents of **2a** and 15 mol % of Fe(acac)<sub>3</sub> were used at 83 °C (SI, Table S2 and S3). Considering that substituents can have a significant influence on the amide formation, a series of isothiureas (**1a-1f**) was prepared *via* our 3CR and tested (SI, Table S4).<sup>10</sup> After all, the substituent on the sulphur and one nitrogen atom can be freely selected by simply altering the thiosulfonate and isocyanide reagents in the activation step. The results indicated that activated aniline equipped with a sterically hindered *N*-*t*-butyl and an *S*-phenyl is the most reactive. When using *N*-*t*-butyl-*S*,*N*-diphenylisothiurea (**1d**) instead of **1a** full conversion can be achieved with only 2.5 mol % catalyst and the desired product **4a** was isolated in 97% (Scheme 2). When the scale of the reaction was increased to 5 mmol (17 times), a similar yield was obtained (SI, Table S4, entry 7).

We first investigated the carboxylic acid scope with activated aniline (Scheme 2). Alkanoic acids (**2**) generally performed very well in reaction with activated aniline **1d**. Branched (**2b**, **2g**), linear (**2c-2e**) as well as cyclic (**2h**) systems gave the desired amides **4** in excellent yields. Also acids featuring a further activation of the methylene  $\alpha$  to the carbonyl by a phenyl group (**2f-g**) selectively reacted with **1d**. Next, we turned our attention to arenecarboxylic acids (**3**) (Scheme 1). Benzoic acid (**3a**) gave *N*-phenylbenzamide (**5a**) in 90% yield.

Electron-donating groups (4-OMe (**3b**), 2-OMe (**3c**), 3,4-Me<sub>2</sub> (**3e**)) and halogens (4-Cl (**3f**), 4-Br (**3g**), 2-I (**3h**)) in different positions of the benzoic acid gave the corresponding products **5b-5h** in 71-88% yields. Even a free hydroxyl group as in 3-hydroxybenzoic acid (**3d**) performed smoothly to provide the corresponding amide **5d** in 85% yield. Notably, also benzoic acids featuring strong electron-withdrawing groups (4-nitro (**3i**), 3-nitro (**3j**), 3,5-bis(trifluoromethyl) (**3k**), 3,4,5-trifluoro (**3l**)) provided the desired products **5i-5l** in 74-95% yield. For most of these benzoic acids, formation of *N-t*-butyl-*N*-phenylurea (**10**) was observed as a side reaction.<sup>14</sup> Substitution of isopropanol by *o*-xylene restored the selectivity, but required a higher reaction temperature. Heteroarene-carboxylic acids such as furan-2-carboxylic acid (**3m**), thiophene-2-carboxylic acid (**3n**), and isomeric pyridine-carboxylic acids (**3o-3q**) also reacted smoothly giving the corresponding products **5m-5q** in good yields (80-99%). Especially the azine type systems (**5o-5q**) are challenging (acids are zwitterions) and converted more slowly. 5 mol% catalyst or a higher boiling alcohol can be used to speed up the reaction in these cases.

**Scheme 2.** Scope of Carboxylic acid **2** and **3**

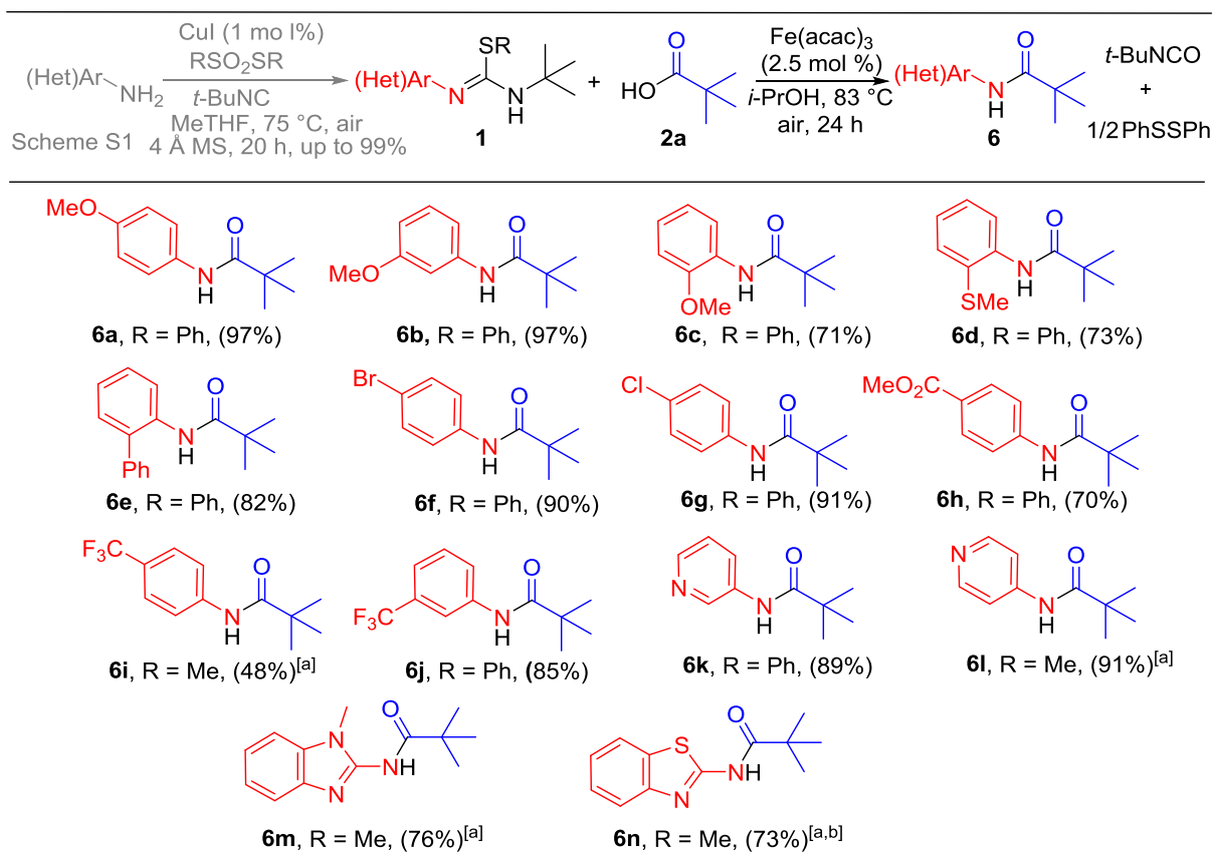


Reaction conditions: **1** (0.3 mmol), **2** or **3** (0.36 mmol), Fe(acac)<sub>3</sub> (2.5 mol %), *i*-PrOH (2 mL), 83 °C, air, 24 h. [a] 72% of PhSSPh was isolated. [b] *o*-xylene (2 mL), 130 °C, 24 h. [c] 2-butanol (2 mL), 98 °C, 48 h. [d] Fe(acac)<sub>3</sub> (5 mol %), 72 h.

The *N*-*t*-butyl-*N'*-(hetero)aryl-*S*-phenylisothiourea (**1**) scope was investigated using pivalic acid (**2a**) as a model sterically encumbered carboxylic acid (Scheme 3). Isothiourea derived from anilines bearing electron-donating substituents (4-OMe (**1h**), 3-OMe (**1i**), 2-OMe (**1j**) and 2-SMe (**1k**)), an *o*-phenyl (**1l**) and halogen substituents (4-Br (**1m**), 4-Cl (**1n**)) underwent the reaction smoothly affording the corresponding products (**6a-6g**) in good to excellent yields. Electron-withdrawing groups (4-CO<sub>2</sub>Me (**1o**), 4-CF<sub>3</sub> (**1p**), 3-CF<sub>3</sub> (**1q**)) were also well tolerated (**6h-j**). Pleasingly, activated heteroareneamines reacted smoothly under standard conditions. Reaction of **2a** with *N*-*t*-butyl-*S*-phenyl-*N'*-(3-pyridinyl)isothiourea (**1r**) gave *N*-(3-pyridinyl)pivalamide (**6k**) in 89%. *N*-*t*-butyl-*N'*-(hetero)aryl-*S*-phenylisothiourea (**1**) containing a (vinylogous) amidine could not be obtained in good yield from the corresponding amine

using our 3CR due to decomposition. Therefore, the corresponding S-methyl analogues were used for these specific heteroareneamines, prepared by using commercial S-methyl methanethiosulfonate as reagent.<sup>10</sup> Both azine (4-aminopyridine (**1s**)) andazole (2-amino-1-methyl-1*H*-benzimidazole (**1t**) and 2-aminobenzothiazole (**1u**)) derived S-methylisothiurea yielded the desired pivalamides in 73-91% yield (**6l-6n**). These are interesting cases as acyl chlorides are known to react with (vinylogous) heteroaromatic amidines at the azine/azole nitrogen first, and therefore consume at least two equivalents of reagent.<sup>15</sup>

**Scheme 3.** Scope of *N*-*t*-Butyl-*N'*-(hetero)aryl-*S*-phenylisothiurea **1**

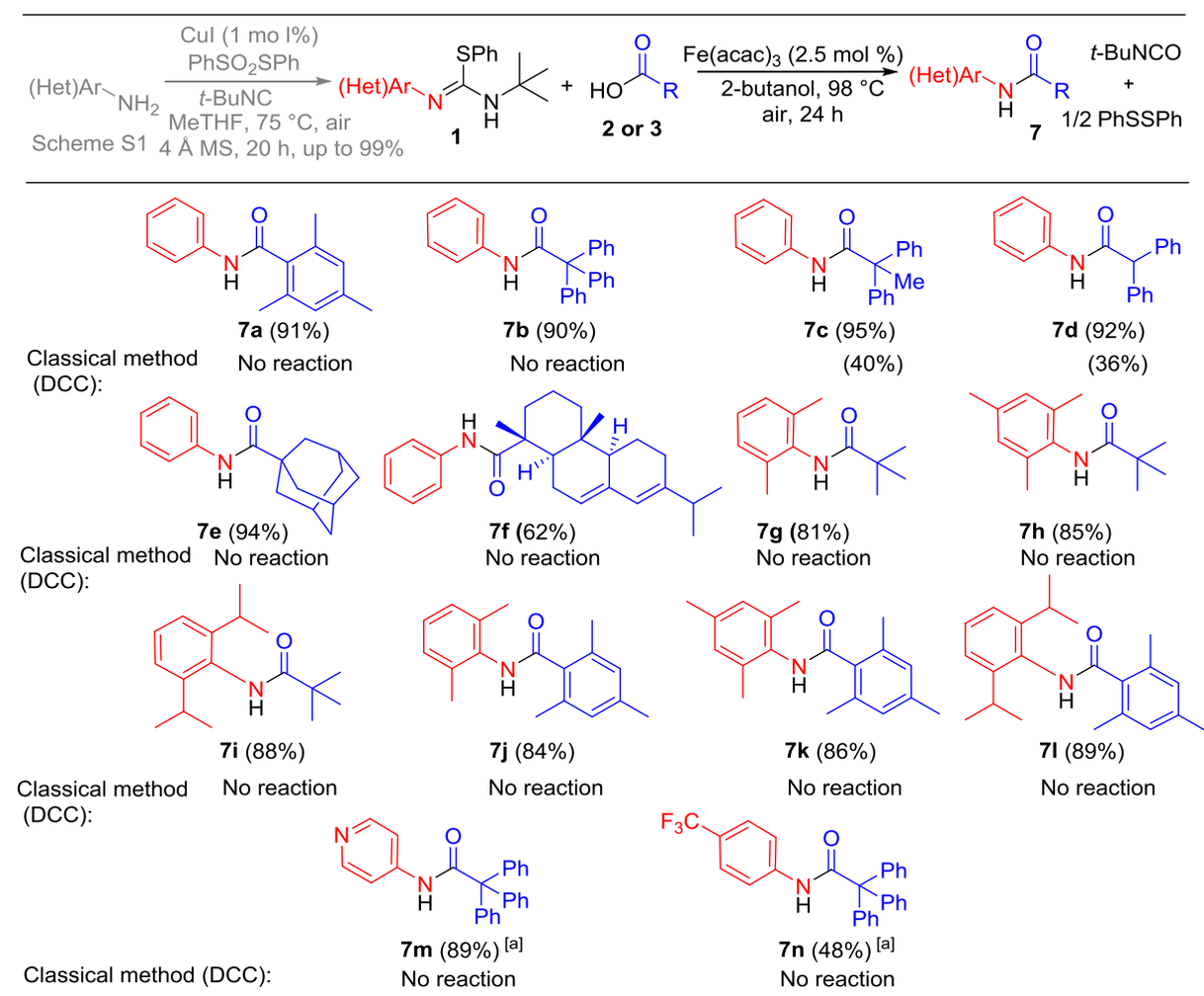


Reaction conditions: **1** (0.3 mmol), **2** or **3** (0.36 mmol),  $\text{Fe}(\text{acac})_3$  (2.5 mol %), *i*-PrOH (2 mL), 83 °C, air, 24 h. [a]  $\text{Fe}(\text{acac})_3$  (15 mol %). [b] *o*-xylene (2 mL), 130 °C, 24 h.

Considering our initial goal, we subsequently investigated the applicability of our protocol for the synthesis of very challenging amides, from the point of view of sterics and electronics. First, reaction of **1d** with carboxylic acids even more sterically hindered than pivalic acid (**2a**) were screened. However, initial tests with 2,4,6-trimethylbenzoic acid (**3r**) revealed the formation of a significant amount of urea **10** under the standard conditions (SI, Table S5). Interestingly, when 2-butanol instead of isopropanol was used as solvent at 98 °C, a selective transformation of **1d**, and 91% of **7a** was achieved (Scheme 4). Under these

conditions sterically hindered aliphatic carboxylic acids (2,2,2-triphenylacetic acid (**2j**), 2,2-diphenylpropanoic acid (**2k**), 2,2-diphenylacetic acid (**2l**), 1-adamantanecarboxylic acid (**2m**) and abietic acid (**2i**)) all reacted with **1d** to provide the corresponding amides **7b-7f** in generally very high yields. Even a combination of a carboxylic acid and an isothiourea which are both sterically hindered was found to be well tolerated in the reaction. Reaction of **2a** or **3r** with *N*-aryl-*N'*-*t*-butyl-*S*-phenylisothiourea featuring a 2,6-Me<sub>2</sub> (**1v**), 2,4,6-Me<sub>3</sub> (**1w**) and 2,6-(isopropyl)<sub>2</sub> (**1x**) phenyl substitution pattern, did not affect the overall efficiency and the corresponding products **7g-7i** were obtained in 81-89%. Also, a combination of a sterically hindered carboxylic acid (**2j**) and an isothiourea derived from an electron deficient (hetero)aromatic amine (4-aminopyridine (**1s**), 4-CF<sub>3</sub> (**1p**)) smoothly give the desired amides **7m** and **7n**.

**Scheme 4.** Synthesis of challenging *N*-(hetero)arylamides **7**



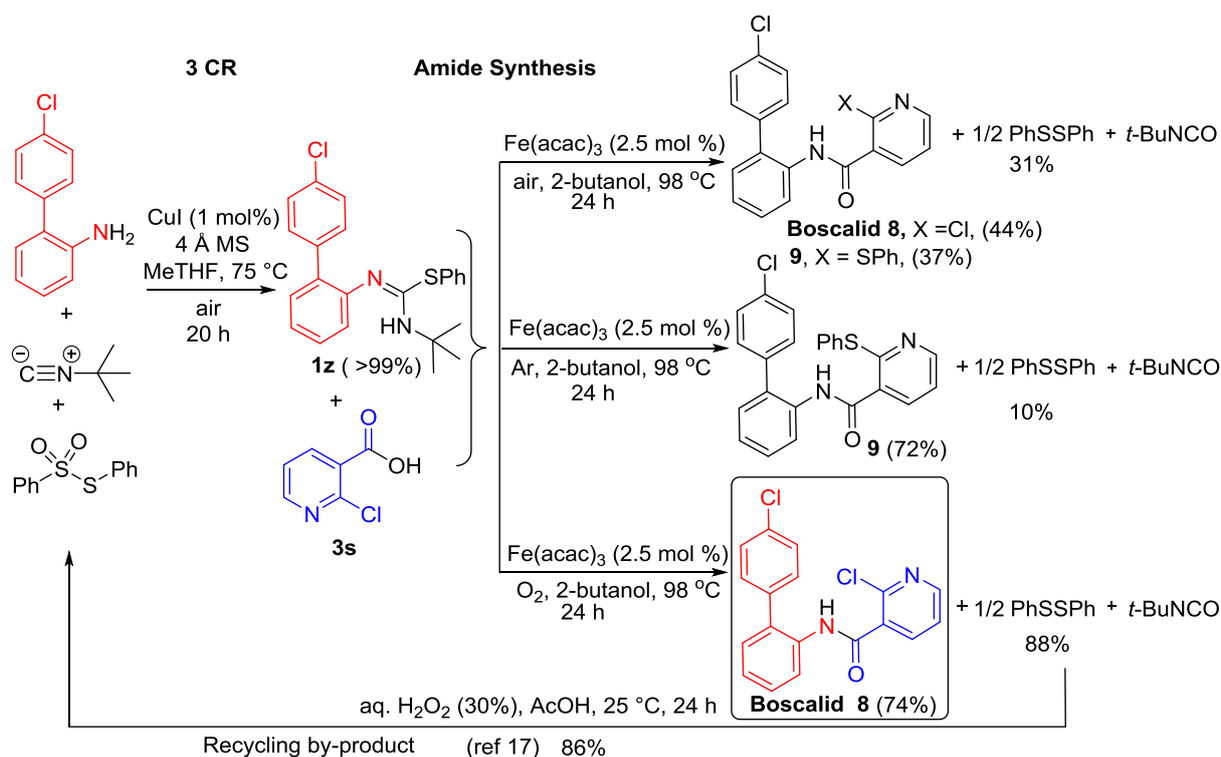
Reaction conditions: **1** (0.3 mmol), **2** or **3** (0.36 mmol), Fe(acac)<sub>3</sub> (2.5 mol %), 2-butanol (2 mL), 98 °C, air, 24 h. [a] *N*-*t*-butyl-*N'*-(hetero)aryl-*S*-methylisothiourea and Fe(acac)<sub>3</sub> (15 mol %). Classical method starting from (Het)ArNH<sub>2</sub>: Carboxylic acid **2** or **3** (1.05 mmol), 1,3-dicyclohexylcarbodiimide (DCC)

(1.08 mmol), 4-dimethylaminopyridine (DMAP) (0.25 mmol), (Het)ArNH<sub>2</sub> (1.0 mmol), CH<sub>2</sub>Cl<sub>2</sub> (10 mL), 0-25 °C, 24 h.

When we attempted to synthesize the challenging amides of Scheme 4 *via* classical reaction of amine and carboxylic acid with DCC no or only a low amount of the target amides was obtained. The new amide synthesis disclosed is therefore complementary with classical protocols. While both protocols will provide standard amides only the new protocol gives efficient access to the challenging representatives.

In order to demonstrate the potential of the new methodology, we applied it to the synthesis of the fungicide Boscalid (Scheme 5).<sup>16</sup> It is classically synthesized by acylation of 2-chloronicotinyl chloride with 4'-chloro-biphenyl-2-ylamine. Our approach started from *N*-*t*-butyl-*N'*-[4'-chloro-(1,1'-biphenyl)-2-yl]-*S*-phenylisothiourea (**1z**) and 2-chloronicotinic acid (**3s**) giving a 74% yield of Boscalid (Scheme 5). **1z** can be prepared in very high yield from commercially available 4'-chloro-biphenyl-2-ylamine, *S*-phenyl benzenethiosulfonate and *t*-butyl isocyanide via the 3CR.<sup>10</sup> It is important that the amide synthesis is in this case executed under oxygen atmosphere, otherwise additional S<sub>N</sub>Ar on the C2-Cl occurs with the thiophenol formed *in-situ*. Under oxygen instead of air the rate of its oxidation to PhSSPh is increased allowing to achieve the desired chemoselectivity. Notably, the PhSSPh by-product formed from PhSH under the amide forming reaction conditions could be isolated in high yield. Interestingly, it can be transformed back into *S*-phenyl benzenethiosulfonate reagent for the 3CR.<sup>17</sup> Also for the model system **4a** (Scheme 2) this recovery procedure was successfully applied (SI, Table S4, entry 7).

**Scheme 5.** Synthesis of Boscalid



In conclusion, we have developed a novel amide synthesis based on rarely explored amine rather than carboxylic acid activation. (Hetero)arylamines are easily activated as bench stable *N*-*t*-butyl-*N'*-(hetero)aryl-*S*-phenylisothiurea via a three component reaction (3CR) with commercially available reagents, *t*-butylisocyanide and *S*-phenyl benzenethiosulfonate. *N*-(hetero)arylamides can be obtained in high selectivity and yield from these isothiureas via reaction with carboxylic acids under iron catalysis, showing a broad functional group compatibility. The protocol allows for the synthesis of very challenging amides, from the point of view of sterics and electronics, not (or only in poor yield) accessible *via* classical coupling agents. The further exploration of the potential of this new methodology as well as the study of its mechanism are currently under investigation in our laboratory.

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## ACKNOWLEDGMENT

This work has been supported by the European Union's Horizon 2020 research and innovation programme under the Marie Skłodowska-Curie grant agreement 657883, the Research Foundation-Flanders (FWO), the UAntwerp (BOF), and the Hercules Foundation. The research leading to these results has also received funding from the Innovative Medicines Initiative ([www.imi.europa.eu](http://www.imi.europa.eu)) Joint Undertaking project CHEM21 under grant

agreement 115360, resources of which are composed of financial contribution from the European Union's Seventh Framework Programme and EFPIA companies' in kind contribution.

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