Ritter-Type Iodo(III)amidation of Unactivated Alkynes for the Stereoselective Synthesis of Multisubstituted Enamides

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Abstract: The Ritter reaction, Brønsted- or Lewis acid-mediated amidation of alkenes or alcohol with nitrile via a carboxylation, represents a classical method for the synthesis of tertiary amides. Although analogous reaction through a vinyl cation or a species alike may offer a route to enamide, an important synthetic building block as well as a common functionality in bioactive compounds, such transformations remain largely elusive. Herein, we report on a Ritter-type transition metal-catalyzed amidation of alkenyl halides, which offers β-iodanyl enamides in moderate to good yields. Mediated by benziodoxole triflate (BXT), the reaction proves applicable to a variety of internal alkynes as well as to various alkyl- and arylalkylamines. The benziodoxole group in the product serves as a versatile handle for further transformations, thus allowing for the preparation of various tri- and tetrasubstituted enamides that are not readily accessible by other means.

Enamides are versatile intermediates for the synthesis of nitrogen-containing molecules.[11] As stable and electron-rich olefins, they participate in diverse transformations such as asymmetric hydrogenation,[2] cycloaddition,[14,2] addition to carbonyls and imines,[3] and halofunctionalization.[4] As for transition metal-catalyzed C–C coupling, enamides have been extensively used in Heck reaction, and more recently have attracted attention as substrates for C–H activation[5] and enantioselective hydroalkynylation.[6] Enamide moiety is also prevalent in bioactive natural products as well as in drug candidates.[11] Conventional methods for the enamine synthesis include condensation of amides and aldehydes, Curtius rearrangement of unsaturated acyl azides, and Wittig or Horner–Wadsworth–Emmons olefination, which often suffer from narrow scope, low stereoccontrol, and/or harsh reaction conditions.[7] More recently, methods based on the functionalization of C≡C bonds have been developed, such as hydroamidation to alkynes[8] and carboamination to ynamides,[9] enabling stereocontrol over the enamide C=C bond.[10] However, the former method is currently limited to terminal alkynes and inherently does not allow access to tetrasubstituted enamides, whereas the latter requires specific types of amide moieties to promote the desired carboamination. The Cu-catalyzed amidation of alkyl halides represents another well-established stereoselective route to enamides,[11] but the synthesis of the prerequisite, stereodefined alkyl halides hinders its application to highly substituted enamides. As such, the chemical space of tetrasubstituted enamides, especially those containing biologically more relevant secondary amide moieties, remains largely inaccessible.[12]

The Ritter reaction, Brønsted- or Lewis acid-mediated reaction of olefins or alcohols with nitrile via carboxylation intermediates, represents a classical but important synthetic approach to tertiary alkyl amides (Scheme 1a).[13] In analogy to the Ritter amidation, one would conceive a possible approach to enamides involving the addition of nitrile to acid- or electrophile-activated alkyne as a vinyl cation-like species, followed by capture of the nitrilium intermediate with water (Scheme 1b). Despite the apparent simplicity, to our knowledge, such transformation on simple unactivated alkynes has remained elusive, while Liu and Yang recently reported a gold-catalyzed Ritter-type amidation of chloro- and bromoalkynes to give trisubstituted, β-haloenamides.[14] Herein, we report that benziodoxole triflate (BXT; 1),[15] a cyclic trivalent iodine electrophile, promotes the Ritter-type difunctionalization of unactivated alkynes to afford tetrasubstituted β-iodanyl enamides in a regio- and stereoselective fashion (Scheme 1c). The reaction tolerates a variety of internal alkynes as well as alkyl- and arylalkylamines. The iodanyl group in the product serves as a handle for the synthesis...
of tri- and tetrastubstituted enamides that are otherwise difficult to access by existing methods.

(a) Ritter reaction: Well studied

(b) Ritter-type functionalization of alkyne: Elusive

(c) This work: Ritter-type iodo(III) amidation of unactivated alkynes

Recently, we reported a trans-iodo(II)etherification reaction of alkynes with BXT and alcohol or with BF₃-activated fluorobenziodoxole (FBX) and dialkyl ether, which allowed for the stereoselective synthesis of multisubstituted vinyl ethers. During this study, we failed in engaging primary and secondary amides as nucleophiles in place of alcohol, observing no desired enamide product. Given this result, we turned our attention to the use of nitrile and water to achieve the putative iodo(III) amidation. In fact, trivalent iodine-mediated functionalization of alkynes with nitrile and water has a precedent. Saito and coworkers reported a synthesis of oxazoles from alkyne, nitrile, and water mediated by iodosylbenzene (PhIO) and a Brønsted acid such as TfOH (Scheme 2a). However, their mechanistic study indicated that this reaction was initiated by the addition of triflate anion, rather than nitrile, to the (I-III)-activated alkyne, and in fact no iodanyl-substituted enamide intermediate was observed. During our study on the iodo(III)etherification, we also observed oxazole formation from 1-phenyl-1-buteny and FBX/BF₃ in wet, HPLC-grade acetonitrile, which was assumed to involve the addition of the nitrile to the BX-activated alkyne (Scheme 2b). Despite these precedent and observation, given the stability of the BX group among other iodanyl groups, we surmised that the desired Ritter-type iodo(III) amidation would be feasible under careful control of water content and other conditions.

Upon extensive screening of reaction conditions (see Tables S1 and S2), we succeeded in the desired iodo(III) amidation of 4-octyne (2a) with 1, acetonitrile, and water under two sets of optimized conditions (Scheme 2c). In conditions A, 2a was reacted with 1 (2 equiv) and H₂O (1 equiv) in the presence of Na₂CO₃ (3 equiv) in dry MeCN (3a) as the solvent at room temperature, affording the desired trans-enamide 4aa in 90% yield with exclusive stereoselectivity. Alternatively, conditions B employed 20 equiv of MeCN in 1,1,1,3,3,3-hexafluoropropanol (HFIP), which also efficiently promoted the reaction to afford 4aa in 88% yield. The yield dropped significantly in trifluoroethanol (TFE), and no desired product was obtained in other solvents such as DCE and toluene. Thus, HFIP appeared to benefit the reaction through its cation-stabilizing nature and low nucleophilicity. The molecular structure of 4aa was unambiguously established by X-ray crystallographic analysis.

Table 1 summarizes the scope of alkynes for the iodo(III)acetamidation using MeCN and H₂O. A variety of dialkylalkynes smoothly participated in the reaction under either conditions A or B to afford the corresponding β-iodenyl enamides 4aa-4ia in moderate to good yields. The preparation of 4aa could be performed on a 2 mmol scale, albeit in somewhat lower yield. For the reactions of unsymmetrically substituted dialkylalkynes, amidation took place preferentially at the sterically more hindered acetylenic carbon, while the degree of steric discrimination was moderate, with regioisomer ratio of ca. 2:1 to 3:1, across alkynes containing methyl group at one end and primary or secondary alkyl group at the other end (see 4ea-4ha). Exclusive regioselectivity was observed in the reaction of 2,2-dimethyloct-3-yn, where the amidation occurred at the carbon proximal to the tert-butyl group (see 4ia and its X-ray structure). Interestingly, this regioselectivity trend marks a sharp contrast with that observed in the iodo(III)etherification of dialkylalkynes with MeOH, where the addition of methoxy group occurred selectively at the sterically less hindered position. This difference between the iodo(III) amidation and the iodo(II)etherification would be ascribed to the sterically less sensitive nature of the nitrile nucleophile compared to the alcohol nucleophile. As such, the reaction would have taken place to avoid the steric repulsion between the BX cation and the alkyl substituent rather than between the nitrile and the alkyn substituent. Besides dialkylalkynes, a series of aryalkylalkynes underwent iodo(III)acetamidation. Thus,
amidation expectedly occurred at the position proximal to the cation-stabilizing aryl group, thus affording the corresponding enamido products 4ja–4pa in moderate yields. As a general trend, conditions A proved suitable for alkynes bearing electron-donating or neutral substituent; they reacted much more sluggishly under conditions B. The opposite was the case for alkynes bearing electron-withdrawing substituents such as ester, ketone, and CF₃ groups. An enyne could also be transformed into dienamide 4qa in 65% yield under conditions A. Note that terminal alkynes failed to give the iodo(III)amidation product under the present conditions.

Table 1. Iodo(III)acetamidation of various alkynes

| Conditions | R' = OAc | R'' = Me | R''' = Me | R'''' = CF₃ | R'''''' = H | R''' = H | Yields |
|------------|----------|---------|---------|------------|-------------|--------|-------|
| A          | 4aa      | 85% (72%) | A       | 87% (B)    |             |        |       |
| B          | 4ba      | 81% (A)  | 88% (B) |             |             |        |       |
| A          | 4ca      | 80% (A)  | 83% (B) |             |             |        |       |
| A          | 4da      | 42% (A)  | 75% (B) |             |             |        |       |
| B          | 4ea      | 74% (A)  | r = 2.3:1 | A           |             |        |       |
| B          | 4fa      | 71% (A)  | r = 2.0:1 | 1.0:1 (B)   |             |        |       |
| A          | 4ga      | 73% (A)  | r = 3.4:1 | A           |             |        |       |
| A          | 4gb      | 74% (A)  | r = 2.2:1 | A           |             |        |       |
| B          | 4gc      | 44% (B)  |        |             |             |        |       |
| A          | 4gd      | 71% (A)  |        |             |             |        |       |
| A          | 4ge      | 44% (B)  |        |             |             |        |       |
| B          | 4gf      | 71% (A)  |        |             |             |        |       |
| A          | 4gg      | 44% (B)  |        |             |             |        |       |
| B          | 4gh      | 71% (A)  |        |             |             |        |       |
| A          | 4gi      | 53% (A)  |        |             |             |        |       |
| B          | 4gj      | 53% (A)  |        |             |             |        |       |

[a] The reaction was performed on a 0.2 mmol scale. The symbol BX in the product formula refers to the benziodoxole moiety. For reactions that produced a mixture of regioisomers, the major isomer is shown along with r (regioisomer ratio) determined by 'H NMR. [b] The yield of a 2-mmol scale reaction (reaction time = 8 h) is given in the parentheses.

We next explored the scope of nitriles in the iodo(III)amidation of 4-octyne (Table 2). Here, conditions B proved effective across a wide variety of organonitriles. Thus, aliphatic nitriles including deuterated acetonitrile ([D₃]-4aa), primary alkyl nitriles (4ab–4ae), benzyl and allyl nitriles (4af–4ah), secondary alkyl nitriles (4ai–4ak), and tertiary alkyl nitriles (4al and 4am) underwent the desired iodo(III)amidation to afford the corresponding β-iodanly enamides in moderate to high yields. The yields were relatively insensitive to the steric size of the nitrile substituent, except that 1-adamantyl nitrile afforded the product 4am in a modest 26% yield. In these reactions, unlike the iodo(III)acetamidation with MeCN, the nitrile substrates were neither dried nor purified prior to the reaction but used as received. Attempted reactions using unpurified propionitrile or butyronitrile as the solvent under conditions A afforded only low yields of the desired enamido, presumably due to the water content in such reagent-grade nitriles. Aromatic nitriles including parent and substituted benzonitriles and thiophene-2-carbonitrile also participated in the reaction with 2a to afford the desired products 4an–4ar in moderate yields.

The product of the present reaction, β-iodanly enamide, provides opportunities for further synthetic transformations via cleavage of the BX group (Scheme 3a) [16a,18d,20,21]. Sonogashira and Stille couplings of 4aa afforded the conjugate enamides 5 and 6, respectively, in good yields. While Sonogashira coupling took place with retention of the olefin stereochemistry, Stille coupling afforded a mixture of E/Z isomers in ca. 5:1 ratio. CuCN/prolinemediated cyanation of 4aa proceeded smoothly at 105 °C to afford the products (E)-7 and (Z)-7 as separable isomers in ca. 3:1 ratio. At a lower temperature of 50 °C, the reaction furnished the β-iodanly enamino 8 with E/Z ratio of 5:1. Hydrodehalogenation of 4aa with Pd/formic acid system afforded the trisubstituted enamino 9 (E/Z = 5:1) in good yield, whereas reductive homocoupling of 4aa mediated by Pd catalyst and Zn afforded the novel diene-diamine 10 (E/Z = 4:1) in excellent yield. It should be pointed out that, unlike the partial loss of the stereochemical integrity in the Pd-catalyzed coupling reactions of 4aa, analogous
reactions of β-iodanyl vinyl ether took place with retention of the stereochemistry. While the reason for this difference remains unclear, we speculate that the vinylpalladium species formed upon oxidative addition of 4aa has a partial carbene-like character due to the electron-donating effect of the amide group, thus allowing rotation of the olefinic C–C bond. The chelating ability of the amide group might also facilitate the isomerization of the vinylpalladium species.

Many of the multisubstituted enamides synthesized above are not readily accessible by other means, and hence would open numerous opportunities for further synthetic transformations. For demonstration, we explored the application to the pyridine/pyrimidine synthesis developed by Movassaghi (Scheme 3b). Thus, trisubstituted enamides obtained through the iodo(III)/amidation/hydrodehalogenation sequence were subjected to TfO, 2-chloropyridine, and electron-rich alkyne or nitrile, affording multisubstituted pyridines 11–14 and pyrimidine 15 in good yields.

In conclusion, we have developed a Ritter-type trans-difunctionalization of alkynes mediated by trivalent iodine electrophile BXT for the synthesis of multisubstituted enamides. The reaction was achieved with careful choice of the reaction conditions including the water content and the reaction medium, and proved applicable to a variety of internal alkynes as well as nitriles, thus affording trans-iodanyl enamides in moderate to good yields. The versatility of the BXT group allows for the preparation of various tri- and tetrasubstituted enamides through cross-coupling. The multisubstituted enamides made accessible by the present method would inspire further development of enamide-based transformations for the synthesis of nitrogen-containing molecules.

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Keywords: enamides • alkynes • amidation • hypervalent iodine compounds • nitriles

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A Ritter-type trans-difunctionalization of alkynes is promoted by benziodoxole triflate, a trivalent iodine electrophile, thus affording tetrasubstituted, β-iodanyl enamides in moderate to good yields. The reaction is applicable to various internal alkynes and nitriles, and transformation of the iodanyl group allows for the preparation of tri- and tetrasubstituted enamides that are difficult to synthesize by other existing methods.