Electrophilic Amide Activation

A Novel Class of 7-Membered Heterocyclic Compounds

Adriano Bauer,[a] Eszter Borsos,[a] and Nuno Maulide*[a]

Abstract: The work presented herein describes the synthesis of a formerly inaccessible class of heterocyclic compounds. The reaction relies on α-phthalimido-amides, which are readily prepared from amino acids in 2 simple reactions steps. Under amide activation conditions in which classical keteniminium ions are not formed, the nitrile solvent is incorporated into the new fused 7-membered ring system. Due to the absence of a keteniminium intermediate, the stereogenic information in the α-position is fully retained.

Introduction

The enhanced nucleophilicity of amides in comparison to other carbonyl functionalities has been noted already in the 19th century.[1] This peculiar reactivity was later leveraged into so-called electrophilic amide activation, a reactivity principle that relies on the treatment of the amide with a strong electrophile such as trifluoromethanesulfonic anhydride (triflic anhydride) and a base in modern approaches (Scheme 1a).[2] In doing so, the amide is transformed into an α-trifloxyenamine 2, which resides in a biased equilibrium with the keteniminium salt 3. The latter, a highly electrophilic species, can be intercepted by several nucleophiles, including nucleophilic oxidants,[3–7] and enables a range of unconventional reactivity profiles for the atoms surrounding the original amide functionality.[8–12] The capture of the central electrophilic atom of the keteniminium by modestly nucleophilic entities such as alkenes (in formal [2+2] cycloaddition reactions[13,14] or other ring-forming events)[15] or ethers[16,17] is a well-documented phenomenon. Although the chemistry of amide activation is dominated by keteniminium formation from tertiary amides, the use of secondary amide starting materials or the outright avoidance of keteniminium species (Scheme 1b)[25] Omission of the base was crucial because it precluded keteniminium formation (and consequently prevented unproductive reaction pathways).

We recently reported that capture of an activated amide, at the iminium triflate stage, by a proximal ester affords previously inaccessible oxazolium salts (Scheme 1b).[25] Omission of the base was crucial because it precluded keteniminium formation (and consequently prevented unproductive reaction pathways).

Herein we would like to report a synthesis of structurally novel 7-membered heterocyclic rings, which similarly relies on the activation of tertiary amid as the absence of keteniminium species (Scheme 1c). This class of compounds has, to the best of our knowledge, not been accessible so far and represents an entirely novel scaffold.

Results and Discussion

In the course of our studies on αβ-dehydrogenation of α-branched amides we investigated the amino-acid derivative 8a under the typical conditions of amide activation (Scheme 2).[4,29] The expected desaturation (to product 10) was observed when the activated amide was treated with pyridine N-oxide in 1,2-DCE upon heating. However, we noticed that when acetonitrile was used as the solvent, the main product of...
the transformation was the 7-membered ring 9a. Inspection of the structure reveals that the nitrile solvent was incorporated into the final product, and indeed nitriles are known to attack activated amides.\[24\] However, in this case, the nitrile-derived fragment is not attached to the former amide functionality. We further noticed that product 9a was not particularly stable and decomposed slowly under air at room temperature upon standing.

During optimization, which was undertaken on substrate 8b, we first noticed that the yield was considerably lower on this very similar substrate under the previously described conditions (Table 1 – Entry 1). Interestingly, the reaction proceeded in the absence of a base (Entry 2/3). Moreover, no desired product was observed when pyridine was used as the base (Entry 4). These results indicated that, similar to our previous study, (Scheme 1b) the avoided formation of a keteniminium intermediate might be crucial to allowing the formation of 9b. Indeed, when the sterically hindered proton scavenger 2,6-ditert-butyl-pyridine (DTBP) was applied, a base that is most likely unable to deprotonate the \(\alpha\)-methine of the activated amide under these conditions, the yield increased considerably (Entry 5). This is consistent with the fact that DTBP was only observed to promote keteniminium formation when less sterically hindered \(\alpha\)-methylene protons were abstracted,\[30\] although the temperature might also play a decisive role.\[31\] Concluding on the experiments of Table 1, the reaction might not need a base that directly participates in the formation of the product. It is more likely that DTBP acts as a proton scavenger to improve the yield. Proton scavengers have been indeed observed to improve the general efficiency of amide activation reactions.\[32\]

With suitable conditions in hand, we investigated if other substrates undergo this unusual reactivity (Scheme 3). First, a range of substituents in the \(\alpha\)-position of the amide was investigated. As shown, variations on the structure of 8a/8b were tolerated. It seems that a number of \(\alpha\)-benzyl or \(\alpha\)-allyl-substituted amides were generally privileged substrates for this transformation (cf. 9d–g). In many cases, the yields determined by \(\text{\textsuperscript{1}H}\) NMR analysis of the crude mixture with mesitylene as the internal standard resulted much higher than the isolated. This is most likely due to the decomposition of the unusual heterocycle during purification via column chromatography. Neither the unsubstituted 9h nor its \(\alpha\)-phenyl-substituted cognate 9i could be generated by this approach, leading instead to complex mixtures. \(\alpha\)-Succinimide amides gave exclusively the starting material back.

For a complete list of references, please see the original paper.
Next, we decided to screen other nitrile solvents and elected to assay butyronitrile as the reaction partner for this transformation (cf. 9j). The resulting product 9j was formed in a good 70 % yield.

Last, we studied the influence of substituents on the phthalimide moiety. While an electron-donating group on the distal position (carbon 11/12) gave a 1:1 mixture of two regioisomers, an electron-withdrawing bromide showed a moderate selectivity for the isomer 9l′. A methyl group in the proximal position (carbon 10/13) showed a rather distinct regioselectivity of 9:1. 2D-NMR analysis of the mixture suggests that the major isomer is surprisingly 9m′, i.e. with the methyl group in direct proximity to the amidine moiety.

The stability of these intriguing structures is, not surprisingly, limited. The remarkable concatenation of an enolester, a 2-azadiene, and a phthalimide-derived amidine is, to the best of our knowledge, unprecedented. Most of the products tended to decompose on silica during column chromatography, systematically resulting in diminished isolated yields when compared to crude NMR assays with an internal standard. When the silica was basified with 1 % of trimethylamine or neutral Al2O3 was used as the stationary phase, products decomposed completely. The use of Florisil® on the other hand, resulted in similar yields (compared to silica gel).

Thereafter mechanistic investigations were undertaken. The reaction mixture becomes strongly colored after triflic anhydride is added and usually turns black after the reaction time of 30 minutes. However, during the aqueous quench, the black turbid color fades quickly to a clear red/orange/yellow solution (depending on the substrate). This observation led us to the assumption that an iminium ion might be transiently formed, which is quickly hydrolyzed to the observed lactone moiety. To support this hypothesis, we quenched the reaction with 18O-labelled water and could indeed observe the full incorporation of the isotope by mass spectrometry (Scheme 4a).

In an additional experiment, the enantioenriched (e.r. > 99 %) phenylalanine derivative (8d) was subjected to the reaction conditions. Interestingly, no racemization of the α-position took place under the optimal reaction conditions (carbon 2 in Scheme 4b), strongly suggesting that this reaction does not proceed via a keteniminium ion. Moreover, we were able to obtain a single crystal of the enantioenriched sample (S)-9d in order to confirm the structure by X-ray analysis.

Concluding from our experiments in Scheme 4a/b we propose the following mechanism for this transformation. Following the formation of O-triflyl oxyiminium ion 10, one of the neighboring phthalimido carbonyl oxygens attacks the highly electrophilic iminium carbon atom. The so generated oxazolinium ion 10 is then attacked by acetonitrile (the solvent) and, upon deprotonation, the ketenimine 12 is generated. This species then undergoes a 5→7 ring enlargement to yield the charged precursor of the final product (14), which is finally converted into the product by aqueous hydrolysis during the workup. The mechanism is somewhat reminiscent of the reversible ring-opening of 4-oxazolines to azomethine ylides which can act as 1,5-dipoles,[33] although 1,3-dipolar cycloadditions are more commonly observed.[34]

Conclusions

In this work, we showed that certain α-phthalimido-amides can be converted into unusual 7-membered heterocyclic systems under electrophilic amide activation in acetonitrile. Noteworthy, our results strongly suggest that the reaction does not proceed via a keteniminium intermediate or analogous vinylic species. The formed products, with a concatenation of sensitive functional groups, can be isolated in yields up to 72 %.
Experimental Section

Reaction Procedure (Scheme 3): The α-phthalimido amide (1.0 equiv., 0.20 mmol) was dissolved in the nitrile (1 mL) and cooled to 0 °C. DTBP was added (2.2 equiv., 0.44 mmol, 99 µL), followed by trifluoromethanesulfonic anhydride (0.22 mmol, 37 µL). The reaction was then stirred at room temperature for 30 minutes. Then, water (2–3 mL) and DCM (2–3 mL) were added to the mixture, resulting in a significant color change. The aqueous phase was extracted with DCM (3 mL) and DCM (2–3 mL) were added to the mixture, resulting in a strong change in color. The aqueous phase was extracted with DCM (2–3 mL) and the united organic layers were dried with Na2SO4 and filtered. Volatiles were removed under reduced pressure and the crude product was purified by column chromatography using silica gel or Florisil® (eluent = heptane/EtOAc, 95:5 → 20:80 v/v%) to obtain the desired product usually as a shiny yellow solid.

Acknowledgments

Funding of this work by the Austrian Science Fund (FWF, P30226) is acknowledged. We thank the University of Vienna for its continued and generous support of our research programs.

Keywords: Amide activation · Cyclization · Heterocycles · 7-Membered rings · Synthetic methods

[1] O. Wallach, Justus Liebigs Ann. Chem. 1877, 184, 1–127.
[2] J.-B. Falmagne, J. Escudero, S. Taleb-Sahraoui, L. Ghosez, Angew. Chem. Int. Ed. Engl. 1981, 20, 879–880; Angew. Chem. 1981, 93, 926.
[3] C. R. Gonçalves, M. Lemmerer, C. J. Teskey, P. Adler, D. Kaiser, B. Maryasin, L. González, N. Maulide, J. Am. Chem. Soc. 2019, 141, 18437–18443.
[4] R. Da Costa, M. Gillard, J. B. Falmagne, L. Ghosez, J. Am. Chem. Soc. 1979, 101, 4381–4383.
[5] D. Kaiser, A. de la Torre, S. Shaaban, N. Maulide, Angew. Chem. Int. Ed. 2017, 56, 5921–5925; Angew. Chem. 2017, 129, 6015.
[6] V. Tona, A. de la Torre, M. Padmanaban, S. Ruider, L. González, N. Maulide, J. Am. Chem. Soc. 2016, 138, 8348–8351.
[7] M. Rens, L. Ghosez, Tetrahedron Lett. 1970, 11, 3765–3768.
[8] D. Kaiser, A. Bauer, M. Lemmerer, N. Maulide, Chem. Soc. Rev. 2018, 47, 7899–7925.
[9] D. Kaiser, N. Maulide, J. Org. Chem. 2016, 81, 4421–4428.
[10] T. Sato, M. Yoritate, H. Tajima, N. Chida, Org. Biomol. Chem. 2018, 16, 3864–3875.
[11] P.-Q. Huang, Acta Chim. Sin. (Engl. Ed.) 2018, 76, 357.
[12] V. Pace, W. Holzer, B. Olofsson, Adv. Synth. Catal. 2014, 356, 3697–3736.
[13] H. Saimoto, C. Houge, A.-M. Hesbain-Frisque, A. Mockel, L. Ghosez, Tetrahydrodron Lett. 1983, 24, 2251–2254.
[14] A. Sidi, J. Marchand-Brynaert, L. Ghosez, Angew. Chem. Int. Ed. Engl. 1974, 13, 267–267; Angew. Chem. 1974, 86, 272.
[15] L. E. Overman, J. P. Wolfe, J. Org. Chem. 2002, 67, 6421–6429.
[16] B. Peng, D. H. O’Donovan, I. D. Jurberg, N. Maulide, Chem. Eur. J. 2012, 18, 16292–16296.
[17] C. Madelaine, V. Valerio, N. Maulide, Angew. Chem. Int. Ed. 2010, 49, 1583–1586; Angew. Chem. 2010, 122, 1628.
[18] P.-Q. Huang, Y.-H. Huang, H. Geng, J.-L. Ye, Sci. Rep. 2016, 6, 28801.
[19] M. Movassaghi, M. D. Hill, O. K. Ahmad, J. Am. Chem. Soc. 2007, 129, 10096–10097.
[20] G. Barbe, A. B. Charette, J. Am. Chem. Soc. 2008, 130, 18–19.
[21] S.-L. Cui, J. Wang, Y.-G. Wang, J. Am. Chem. Soc. 2008, 130, 13526–13527.
[22] W. S. Bechara, G. Pelletier, A. B. Charette, Nat. Chem. 2012, 4, 228–234.
[23] O. K. Ahmad, J. W. Medley, A. Coste, M. Movassaghi, Org. Synth. 2012, 89, 549.
[24] M. Movassaghi, M. D. Hill, J. Am. Chem. Soc. 2006, 128, 14254–14255.
[25] E. Spinozzi, A. Bauer, N. Maulide, Eur. J. Org. Chem. 2019, 5230–5233.
[26] A. B. Charette, P. Chua, J. Org. Chem. 1998, 63, 908–909.
[27] P. Cyr, S. Régnier, W. S. Bechara, A. B. Charette, Org. Lett. 2015, 17, 3386–3389.
[28] W. S. Bechara, I. S. Khazhieva, E. Rodriguez, A. B. Charette, Org. Lett. 2015, 17, 1184–1187.
[29] A. Bauer, N. Maulide, Chem. Sci. 2019, 10, 9836–9840.
[30] L. Chen, L. Ghosez, Tetrahedron Lett. 1990, 31, 4467–4470.
[31] H. Chen, Y.-H. Huang, J.-L. Ye, P.-Q. Huang, J. Org. Chem. 2019, 84, 9270–9281.
[32] B. Mátrovölgyi, T. Hergert, E. Bálint, P. Bagi, F. Faigl, J. Org. Chem. 2018, 83, 2282–2292.
[33] T. Hiyama, H. Taguchi, H. Nozaki, Bull. Chem. Soc. Jpn. 1974, 47, 2909–2910.
[34] E. Vedejs, J. W. Grissom, J. Org. Chem. 1988, 53, 1876–1882.