

Loyola University Chicago

Loyola eCommons

Chemistry: Faculty Publications and Other Works

Faculty Publications and Other Works by Department

8-2015

Metal-Free Tandem Beckmann-Electrophilic Aromatic Substitution Cascade Affording Diaryl Imines, Ketones, Amines, and Quinazolines

Samuel Sarsah Loyola University Chicago, samuelsarsah@yahoo.com

Marlon R. Lutz Jr. Loyola University Chicago

Kailyn Chichi Bobb Loyola University Chicago

Daniel Becker Loyola University Chicago, dbecke3@luc.edu

Follow this and additional works at: https://ecommons.luc.edu/chemistry_facpubs



Part of the Biochemistry Commons, and the Chemistry Commons

Author Manuscript

This is a pre-publication author manuscript of the final, published article.

Recommended Citation

Sarsah, Samuel; Lutz, Marlon R. Jr.; Bobb, Kailyn Chichi; and Becker, Daniel. Metal-Free Tandem Beckmann-Electrophilic Aromatic Substitution Cascade Affording Diaryl Imines, Ketones, Amines, and Quinazolines. Tetrahedron Letters, 56, 40: 5390-5392, 2015. Retrieved from Loyola eCommons, Chemistry: Faculty Publications and Other Works, http://dx.doi.org/10.1016/j.tetlet.2015.07.095

This Article is brought to you for free and open access by the Faculty Publications and Other Works by Department at Loyola eCommons. It has been accepted for inclusion in Chemistry: Faculty Publications and Other Works by an authorized administrator of Loyola eCommons. For more information, please contact ecommons@luc.edu.



This work is licensed under a Creative Commons Attribution-Noncommercial-No Derivative Works 3.0 License. © Elsevier Ltd. 2015

Metal-Free Tandem Beckmann-Electrophilic Aromatic Substitution Cascade Affording Diaryl Imines, Ketones, Amines and Quinazolines

Tetrahedron Lett **2015** *56* 5390–5392

Samuel R.S. Sarsah, Marlon R. Lutz, Jr., Kailyn Chichi Bobb, and Daniel P. Becker*

Metal-Free Tandem Beckmann-Electrophilic Aromatic Substitution Cascade Affording Diaryl Imines, Ketones, Amines and Quinazolines

Samuel R.S. Sarsah[‡], Marlon R. Lutz[§], Jr., Kailyn Chichi Liu, Daniel P. Becker*

Department of Chemistry, Loyola University Chicago, 1032 West Sheridan Road, Chicago, IL 60660, dbecke3@luc.edu

‡Current Address: Research & Development, Star Thermoplastic Alloys and Rubbers Inc., 2121 W 21st St, Broadview, IL 60155, ssarsah@luc.edu
§Current Address: Process Research and Development, Regis Technologies Inc., 8210 Austin Avenue, Morton Grove, IL 60053, mlutz@registech.com

Abstract

A cascade reaction sequence involving a Beckmann rearrangement on benzophenone oxime followed by an electrophilic aromatic substitution (EAS) on the intermediate nitrilium ion affords N-phenyl diaryl imines that may then be hydrolyzed to ketones, or reduced to the corresponding amines. Reaction with benzonitrile afforded 2,4-diphenylquinazoline through a Beckmann-Ritter-EAS cascade.

Cascade or domino reactions enable the assemblage of complex molecules through a sequence of reactions where one reaction step prepares a reactive intermediate that is immediately employed in a subsequent reaction.¹ Examples of cascade reactions include the tandem Aldol-Tishchenko reaction^{2,3} and the Banert cascade.⁴ The Beckmann rearrangement⁵ is a synthetic workhorse that is still of current theoretical interest regarding the concerted or stepwise nature of its mechanism.⁶ As part of our program to investigate apex-modified derivatives⁷ of the bowl-shaped supramolecular scaffold cyclotriveratrylene (CTV),⁸ we employed the Beckmann rearrangement to afford a ring expansion via the oxime 1⁹ derived from CTV to afford the corresponding 10-membered CTV lactam 3, which depending upon the experimental conditions, was also accompanied by the product of an intramolecular electrophilic addition reaction

affording an unexpected helical pentacycle 4 (Scheme 1). We realized that this byproduct must have arisen from a cascade process wherein the intermediate nitrilium ion 2 formed in the Beckmann reaction was intercepted by one of the electron-rich veratrole rings of the macrocycle in a trans-annular process. We questioned whether this tandem Beckmann-electrophilic aromatic addition reaction could be generalized to an intermolecular variant with electrophilic aromatic substitution arising from the Beckmann intermediate, and were encouraged by similar cascade processes. Schinzer previously employed nucleophilic allylsilanes to intramolecularly trap the cationic Beckmann rearrangement intermediate in the preparation of various heterocycles. 11-13 Amidines have been synthesized by trapping the Beckmann iminocarbocation intermediates derived from oximes by Katritzky¹⁴ while Mukaiyama conveniently formed amidines and enamines by trapping Beckmann carbocation intermediates with amines and resonancestabilized carbanion nucleophiles, respectively using trifluoromethanesulfonic anhydride as the electrophilic initiator, 15 but trapping of the cationic Beckmann rearrangement intermediate by an intermolecular electrophilic aromatic addition has not been previously reported to our knowledge. We describe herein the intermolecular tandem Beckmannelectrophilic aromatic substitution cascade utilizing benzophenone oxime and a variety of aromatic nucleophiles to afford the corresponding imines which could be isolated, hydrolyzed, or reduced to the corresponding amines.

Scheme 1: Intramolecular Tandem Beckman-Electrophilic Aromatic Addition from CTV Oxime

In the original intramolecular case, the reaction is favored both by the very electron-rich nature and close proximity (3.4-3.6Å)¹⁰ of the dimethoxy ring to the Beckmann nitrilium intermediate. We therefore initially employed neat conditions and electron rich π nucleophiles to probe the viability of the intermolecular variation, employing benzophenone oxime as the Beckmann substrate. Initiating the Beckmann with thionyl chloride did not afford the tandem product, as it led only to the formation of the simple Beckmann product, benzanilide, as rapid attack of the chloride anion precluded attack by the π -nucleophile. Thus, an intermediate chloroimine presumably formed which was hydrolyzed upon workup. Switching to trifluoromethanesulfonic anhydride to initiate the Beckmann rearrangement¹⁵ in the absence of a nucleophilic leaving group allowed time for the nitrilium ion to react with the π -nucleophilic aromatic to give good to excellent yields of the TB-EAS products (Table 1). In the first case with the π -nucleophile veratrole employed as solvent, a 95% isolated yield of the hydrolyzed ketone 6a was obtained. We continued to explore the generality of the TB-EAS sequence employing various electron rich and electron poor π -nucleophiles, first under neat conditions, followed by hydrolyzing the intermediate imine 7 directly to the ketone products (6). We examined reducing the ratio of π -nucleophile to two equivalents relative to benzophenone oxime in 1,2-dichloroethane at reflux (Table 2). Once validated through isolation of the ketone, the imines 7 were isolated, and the subsequent imines were reduced to the corresponding α , N-arylbenzenemethanamines **8**, as these molecules as a class are important in pharmaceuticals¹⁶.

Scheme 2: Preparation of diarylketones 6 and amines 8 via imines 7

Table 1. TB-EAS Results with π -Nucleophiles^a

Entry	π-Nuc	Ar	Ketone 6 ^b Yield (% neat/soln)	Imine 7° Yield (%)	Amine 8 Yield (%)
a	veratrole	3,4-dimethoxyphenyl	95/91	58 ^f	51 ^d
ь	1,4- dimethoxy- benzene	2,5-dimethoxyphenyl	97/72	56	57 ^d
С	anisole	4-methoxyphenyl (plus ortho)	95/96	91 ^f	90°
d	<i>p</i> -xylene	2,5-dimethyl phenyl	95/70	90	63 ^e
e	PhCH ₃	4-methylphenyl (plus ortho)	75/93	66 ^f	73°
f	PhCl	4-chlorophenyl (plus ortho)	68/19	56 ^f	45°
g	PhBr	4-bromophenyl (plus ortho)	75/34	60 ^f	59 ^e
h	PhCN	3-cyanophenyl	$0^{\mathrm{g}}/0^{\mathrm{g}}$	39 ^g	

^aBenzophenone oxime in 1,2-dichloroethane was treated with triflic anhydride under reflux for 16-24h in the presence of 2 equivalents of π-nucleophile with the exception for PhCl and PhBr which used 4 eq and 8 eq, respectively. ^bHydrolytic workup with acid afforded ketones **6**; ^c aqueous sodium bicarbonate quench afforded imines **7**; ^dCrude imines were treated with NaBH₄ in MeOH or THF to form amines **8**; ^eReflux with lithium aluminum hydride in THF/2h followed by Fieser workup afforded amines **8**; ^fThe imine contained traces of ketone after workup; ^gReaction with benzonitrile gave the

imine even after acidic hydrolytic workup, along with 2,4-diphenylquinazoline (**10** in 47% yield).

Encouraged by the success with neat veratrole, but wishing to avoid the inconvenience of using neat π -nucleophiles as solvent, we obtained a 91% yield of ketone **6a** employing 2 equivalents of veratrole in 1,2-dichloroethane. In a separate experiment, veratrole afforded a 58% yield of imine 7a after column chromatography, generally as a mixture of E and Z isomers, and contaminated in most cases with some ketone 6a. The imines were difficult to isolate as they suffered hydrolysis during purification, and were best reduced directly. Direct reduction of the imine 7a after the TB-EAS sequence furnished a 51% yield of α,N-arylbenzenemethanamine 8a (R=H). Ketone products 6a-g could also be obtained by acid hydrolysis of the isolated imine, but as expected, the yields were lower than the direct hydrolysis. Amine products (8a-g) were obtained by direct reduction of the crude imines from the TB-EAS sequence without isolation. Employing pdimethoxybenzene gave a high yield of ketone **6b** (97%) after hydrolysis but attempted isolation of the imine was problematic due to very rapid hydrolysis even with an aqueous bicarbonate workup. Direct reduction of the imine with sodium borohydride gave amine **8b** in 57% yield. Neat anisole provided a 95% yield of ketone **6c** (7:1 para:ortho) and reaction with 2 equivalents of anisole in 1,2-dichloroethane at reflux gave a comparable high isolated yield of ketone 6c (96%). A mild basic quench enabled isolation of imine 7c in 91% yield. Direct reduction of the imine to α , N-(4-methoxyphenyl)benzenemethanamine with sodium borohydride proceeded in 90% yield. The sigma donor, p-xylene, furnished a 95% isolated yield of diaryl ketone 6d employing p-xylene as the solvent, although the use of 2 equivalents of *p*-xylene resulted in a moderately lower yield (70%). The imine 7d derived from p-xylene was isolated in 90% yield, whereas reduction of imine 7d gave amine 8d in 63% yield. Performing the TB-EAS reaction with toluene afforded a more modest yield under neat conditions (75%) but surprisingly a higher yield (93%) when using 2 equivalents of toluene in 1,2dichloroethane. The imine 7e from toluene was isolated in 66% yield, and was reduced to aryl amine **8e** in 73% yield. The TB-EAS reactions with halobenzenes were more sluggish and gave lower yields. Chlorobenzene and bromobenzene gave ketones 6f and

6g in moderate yields (68% and 75%, respectively), but yields were lower in solution, despite employing 4 and 8 equivalents of the halobenzene, respectively (19% and 34%), whereas the 4-chlorophenylimine **7f** and 4-bromophenylimine **7g** intermediates were isolated in moderate yields (56% and 60%, respectively). Thus, the bromo and chlorobenzene nucleophiles were considerably more concentration dependent and relatively higher temperatures are needed to achieve moderate yields; reaction at 2M concentration of nucleophile proved to be unsuccessful whereas reaction at 4M produced low yields and 8M gave the yields shown. The amines **8f** and **8g** were isolated in 45 and 59% yield, respectively. Utilization of nitrobenzene as the π -nucleophile (not shown) gave low yields of as-yet unidentified products and was not pursued.

Employing neat benzonitrile as the π -nucleophile in the TB-EAS sequence did not give the expected ketone 6h upon acidic hydrolysis, but rather imine 7h was isolated in 39% yield, along with a 47% yield of 2,4-diphenylquinazoline 10a (Scheme 3), which arises from a Ritter type reaction 17-18 involving attack of the Beckmann nitrilium ion by benzonitrile followed by an EAS reaction by the N-phenyl group on the resulting nitrilium ion 9a/b, comprising a Beckmann-Ritter-EAS cascade sequence. Ritter iminium ions have been trapped previously by intramolecular EAS to afford dihydroisoguinolines, ¹⁹ and Kofanov reported a similar cascade sequence starting with an N-aryl amide to prepare 2,4-diarylquinazolines.²⁰ The intermediate in this sequence should be the same cation proposed by Meerwein in his early synthesis of quinazolines where he treated the corresponding chloroiminium species with aluminum trichloride.²¹ When we employed benzonitrile in solution (8M in 1,2-dichloroethane) rather than in the neat reaction, a much improved (88%) yield of 2.4-diphenylquinazoline 10a (R = Ph) was obtained. On the other hand, employing acetonitrile as the solvent at 75°C afforded 2phenyl-4-methylquinazoline 10b ($R = CH_3$) in 37% isolated yield, but acetonitrile as a solution in 1,2-dichloroethane afforded 10b in only 13% yield). Thus, the Beckmann-Ritter-EAS cascade affording quinazolines is higher yielding with the aryl nitrile versus the aliphatic nitrile under the present conditions, but the generality with more substituted derivatives in both categories remains to be explored.

Scheme 3: Preparation of quinazolines 10a (R = Ph) and 10b (R = CH₃)

In summary, the intermolecular TB-EAS reaction commencing with benzophenone oxime affords N,α,α-triarylimine derivatives which may be hydrolyzed to unsymmetrical diarylketones or reduced to α,N-arylbenzenemethanamines. Reaction with benzonitrile proceeded via a Beckmann-Ritter-EAS cascade sequence to afford 2,4-diphenylquinazoline (10a) in very good (88%) yield. Current efforts are focused on expanding the scope of the TB-EAS and Beckmann-Ritter-EAS cascade sequences, particularly toward the synthesis of more highly substituted quinazolines under metal-free conditions.

Acknowledgments

Loyola University Chicago is gratefully acknowledged for support of this research and for a Mulcahy Scholarship for K.C.B under the auspices of LUROP (Loyola Undergraduate Research Opportunities Program).

Supplementary data

Supplementary data associated with this article can be found in the online version.

References

- (1) Jones, S. B.; Simmons, B.; Mastracchio, A.; MacMillan, D. W. C. *Nature (London, U. K.)* **2011**, *475*, 183-188.
- (2) Bodnar, P. M.; Shaw, J. T.; Woerpel, K. A. J. Org. Chem. 1997, 62, 5674-5675.
- (3) Honda, M.; Iwamoto, R.; Nogami, Y.; Segi, M. Chem. Lett. 2005, 34, 466-467.
- (4) Loren, J. C.; Sharpless, K. B. Synthesis **2005**, 1514-1520.
- (5) Gawley, R. E. Organic Reactions (Hoboken, NJ, United States) 1988, 35, 1-420.
- (6) Yamabe, S.; Tsuchida, N.; Yamazaki, S. J. Org. Chem. 2005, 70, 10638-10644.
- (7) Panagopoulos, A. M.; Zeller, M.; Becker, D. P. J. Org. Chem. 2010, 75, 7887-7892.
- (8) Collet, A. Tetrahedron 1987, 43, 5725-5759.

- (9) Lutz Jr., M. R.; French, D. C.; Rehage, P.; Becker, D. P. *Tetrahedron Letters* **2007**, 48, 6368-6371.
- (10) Lutz, M. R., Jr.; Zeller, M.; Becker, D. P. Tetrahedron Lett. 2008, 49, 5003-5005.
- (11) Schinzer, D.; Bo, Y. Angewandte Chemie Chem., Int. Ed. Engl., 1991, 30, 687-8.
- (12) Schinzer, D.; Langkopf, E. Synlett **1994**, 375-377.
- (13) Schinzer, D.; Abel, U.; Jones, P. G. Synlett 1997, 632-634.
- (14) Katritzky, A. R.; Monteux, D. A.; Tymoshenko, D. O. Org. Lett. 1999, 1, 577-578.
- (15) Takuwa, T.; Minowa, T.; Onishi, J. Y.; Mukaiyama, T. Chem. Lett. 2004, 33, 322-323.
- (16) Lu, Y.; Nikolovska-Coleska, Z.; Fang, X.; Gao, W.; Shangary, S.; Qiu, S.; Qin, D.; Wang, S. *J. Med. Chem.* **2006**, *49*, 3759-3762.
- (17) Ritter, J. J.; Minieri, P. P. J. Am. Chem. Soc. 1948, 70, 4045-4048.
- (18) Krimen, L. I.; Cota, D. J. Org. React. 1969, 17, 213-325.
- (19) Janin, Y. L.; Decaudin, D.; Monneret, C.; and Poupon, M.-F. Tet. **2004** *60* 5481-5485.
- (20) Kofanov, E. R.; Sosnina, V. V.; Danilova, A. S.; Korolev, P. V. Russian Journal of Applied Chemistry 1999, 72, 850-852.
- (21) Meerwein, H.; Laasch, P.; Mersch, R.; Nentwig, J. Chem Ber. 1956, 89, 224-238.