Tetrahedron Letters 50 (2009) 75-76

Contents lists available at ScienceDirect

Tetrahedron Letters

journal homepage: www.elsevier.com/locate/tetlet



Synthesis of 4-, 5-, 6-, and 7-azidotryptamines

Anne Friedrich^{a,b}, Stefan Bräse^b, Sarah E. O'Connor^{a,*}

^a Massachusetts Institute of Technology, Department of Chemistry, 77 Massachusetts Ave., Cambridge, MA 02139, USA
^b Institut f
ür Organische Chemie, Universit
ät Karlsruhe (TH), Fritz-Haber-Weg 6, D-76131 Karlsruhe, Germany

ARTICLE INFO

Article history:

Received 12 June 2008

Revised 18 October 2008

Accepted 20 October 2008 Available online 1 November 2008

ABSTRACT

Synthesis of azidotryptamines from commercially available nitroindoles via the corresponding amino tryptamines in good overall yields (15–38%) is presented.

© 2008 Elsevier Ltd. All rights reserved.

Photoaffinity-derivatized ligands have been used to covalently label and to identify numerous classes of enzymes. Aryl azides, diazirines, and benzophenones have all been used as photolabels.¹ The use of azidoindoles for photocrosslinking is precedented; for example, 6-azidotryptophan has been used to label the tryptophan $\alpha_2\beta_2$ synthase complex,² and 5-azidoindole-3 acetic acid has been shown to efficiently cross-link to auxin receptors in plant cell extracts.³ Despite the utility of the indole azide labeling group, and the participation of tryptamine in metabolic pathways, no synthetic route had been previously reported for the azidotryptamines **1**. Herein, we report an efficient synthesis of 4-, 5-, 6-, and 7-azidotryptamines **1a–d** starting from commercially available nitroindoles. Adapting a previously reported protocol for tryptamine synthesis, nitroindoles **2a,b,d** were reacted with oxalylchloride in ether (Scheme 1A).⁴ The resulting indole oxalylchlorides were treated with ammonia in 1,4-dioxane to give the corresponding oxalylamides **3a,b,d** in good yields. Simultaneous reduction of the nitro group and amide side chain with lithium aluminum hydride in refluxing THF/1,4-dioxane led to the aminoindoles **4a,b,d** (Scheme 1A). Surprisingly, 6-nitroindole **2c** failed to react with oxalylchloride to form the desired product under a variety of temperatures and reagent concentrations. Therefore, an alternative strategy for the synthesis of the 6-aminoindole **4c** was employed (Scheme 1B). Nitroindole-3-carboxaldehyde **5c** was prepared by utilizing the *Vilsmeier–Haack* reaction, followed by addition of nitromethane



Scheme 1. (A) Synthesis of 4-, 5-, and 7-aminotryptamine 4a,b,d from nitroindoles 2a,b,d via the indole oxalylamides 3a,b,d. (B) Synthesis of 4-, 5-, and 7-aminotryptamine 4b,c,d from nitroindoles 2b,c,d via the nitro vinyl indoles 6b,c,d.

^{*} Corresponding author. Tel.: +1 617 324 0180; fax: +1 617 324 0505. *E-mail address*: soc@mit.edu (S. E. O'Connor).

^{0040-4039/\$ -} see front matter @ 2008 Elsevier Ltd. All rights reserved. doi:10.1016/j.tetlet.2008.10.091



Scheme 2. Diazotation of aminotryptamines 4a-d to form azidotryptamines 1a-d.







Scheme 3. Chemical reaction of tryptamine **1a-d** and secologanin **8** to yield strictosidine **7a-d**, which appears to be incorporated into several terpene indole alkaloids including **9**, **10**, and **11**.

and elimination (*Henry* reaction) to give nitro vinyl indole **6c**. Reduction with lithium aluminum hydride in refluxing THF gave 6-amino-tryptamine **4c**. This method could also be used to generate 5-aminotryptamine **2b** and 7-aminotryptamine **2d**, though addition/elimination and reduction of **2a** to **4a** did not proceed in good yields through this route (Scheme 1B).⁵

With aminotryptamines **4a–d** in hand, the azide group could be introduced by diazotation of the corresponding amines. Following a protocol from Melhardo et al., sodium nitrite and sodium azide in glacial acetic acid were used to convert the aminotryptamines **4a–d** into the azidotryptamines **1a–d**.⁶ The primary alkyl amine did not react under these conditions, and all four azidotryptamines **1a–d** (Scheme 2). From the nitroindoles **2a–d**, the corresponding azidotryptamines **1a–d** could be obtained in overall yields ranging from 15% to 38%.

A concern when using photoaffinity-derivatized substrates to identify proteins is that the azide group could disrupt binding to highly substrate-specific enzymes. To examine this issue, we chemically synthesized the metabolic biosynthetic intermediate strictosidine **7a–d** from **1a–d** (Scheme 3). Photolabeled azidostrictosidines **7a–d** were incubated with *Catharanthus roseus* plant cell culture that produces monoterpene indole alkaloids ajmalicine (m/z 353) **9**, serpentine (m/z 349) **10**, and tabersonine (m/z 337) **11** that are derived from strictosidine **7**.⁷ Mass spectrometry analysis of these *C. roseus* extracts revealed the formation of new compounds displaying masses consistent with azido analogs of alkaloids with m/z 353.⁸ Furthermore, in cultures supplemented with azidostrictosidines **7c** and **7d**, compounds with molecular formulae consistent with azido analogs of alkaloids having m/z 349⁹ and m/z 337¹⁰ were also observed. These compounds were not observed in control cultures lacking azidostrictosidine.

Although MS analysis cannot allow us to predict the structure of these unknown analogs, these studies nevertheless strongly suggest that the biosynthetic enzymes of an alkaloid metabolic pathway can bind to and turn over azide-labeled precursors. Biosynthetic intermediates derived from 1a-d and 7a-d may therefore potentially be used for photoaffinity labeling of enzymes in this metabolic pathway. In combination with a chemoselective handle (such as an alkyne installed at the ester of 7)¹¹ that allows for identification in a crude mixture, these photolabeled compounds could be used to identify desired metabolic enzymes in cell lysates.

Acknowledgments

We gratefully acknowledge support from the Koch Fund and GM074820, as well as support from Landesgraduiertenförderung Baden-Württemberg and DAAD for fellowship support for A.F. We thank Elizabeth McCoy for helpful discussion.

Supplementary data

Supplementary data (experimental protocols and spectroscopic characterization of compounds **1–5a–d**, **7a–d**) associated with this article can be found, in the online version, at doi:10.1016/j.tetlet.2008.10.091.

References and notes

- 1. Dorman, G. Top. Curr. Chem. 2001, 211, 169–225.
- 2. Miles, E. W.; Phillips, R. S. Biochemistry 1985, 24, 4694-4703.
- 3. Zettl, R.; Schell, J.; Palme, K. Proc. Natl. Acad. Sci. U.S.A. 1994, 91, 689-693.
- 4. Wu, T. Y. H.; Schultz, P. G. Org. Lett. 2002, 4, 4033-4036.
- An older report of the synthesis of aminotryptamines is described in: Hiremath, S. P.; Siddappa, S. J. Med. Chem. 1965, 8, 142–143. However, no yields or spectroscopic characterization were provided in this study.
- (a) Li, M.; Johnson, M. E. *Tetrahedron Lett.* **1994**, 35, 6255–6258; (b) Melhardo, L. L.; Leonard, N. J. *J. Org. Chem.* **1983**, 48, 5130–5133.
- (a) Hamill, J. D.; Parr, A. J.; Rhodes, M. J. C.; Robins, R. J.; Walton, N. J. Bio/ Technology **1987**, 5, 800–804; (b) Rijhwani, S. K.; Shanks, J. V. Enzyme Microb. Technol. **1998**, 22, 606–611.
- Azido analog of [M+H]⁺ 353 (e.g., 9). Expected [M+H]⁺ 394.1879. Observed [M+H]⁺ after co-culture with: *no substrate* not observed; 7a 394.1886; 7b 394.1884; 7c 394.1873; 7d 394.1893.
- Azido analog of [M]⁺ 349 (e.g., 10). Expected [M]⁺ 390.1566. Observed [M+H]⁺ after co-culture with: *no substrate* not observed; 7a not observed; 7b not observed; 7c 390.1581; 7d 390.1581.
- Azido analog of [M+H]⁺ 337 (e.g. 11). Expected [M+H]⁺ 378.1930. Observed [M+H]⁺ after co-culture with: *no substrate* not observed; 7a not observed; 7b not observed; 7c 378.1933; 7d 378.1943.
- 11. Galan, M. C.; McCoy, E.; O'Connor, S. E. Chem. Commun. 2007, 3249-3251.