L-Proline: An Efficient and Selective Catalyst for Transamidation of Thioamides with Amines

Sadu Nageswara Rao, Darapaneni Chandra Mohan and Subbarayappa Adimurthy*
Academy of Scientific and Innovative Research, CSIR–Central Salt and Marine Chemicals Research Institute, GB Marg, Bhavnagar-364002, Gujarat, India

Abstract
L-Proline catalysed transthioamidation of primary thioamides with amines under solvent-free conditions has been described. The transthioamidation is compatible with a wide range of amines with yields up to 97%.

Keywords: Thioamides; Amines; L-Proline; Transthioamidation

Introduction
Thioamide is an important and useful functional group in both chemistry and biology. Thioamides are not only serve as versatile synthetic intermediates for the construction of pharmaceutically important molecules containing nitrogen and sulfur heterocycles [1-7] but are also used as antitumor agents and enzyme inhibitors [8-10]. Thioamide-based drugs such as ethionamide (ETH) and prothionamide (PTH) have been widely used for many years in the treatment of mycobacterial infections caused by Mycobacterium tuberculosis, M. leprae and M. avium complex infections [11,12]. Recently, functionalized-thioamide fluorescent dyes were also employed as metal ion sensors [13]. Diverse synthetic methods have been discovered for the synthesis of thioamides [14-23].

Transamidation is an attractive tool represents one of the most convenient and straightforward method, that would exchange the constituents of two different amide groups. Compared with transamidation of amides with amines, the corresponding transthioamidations are rarely reported with rather limited substrate scope [24,25].

Most of the approaches for thioamide syntheses require transition metal catalysts to promote this transformation efficiently; also they suffer from inadequacies such as the expensive nature of catalyst, moisture and/or air sensitivity of Grignard reagents. Thus, the separation of metal catalyst from products, which is of particular importance for the synthesis of pharmaceuticals and fine chemicals because of their residual toxicity in the target compounds, is a central issue to consider. Moreover, transition metal-catalysed reactions also generate hazardous waste which is environmentally problematic and hence, should be avoided wherever possible. These catalysts are active only in organic solvents. Therefore, the development of transthioamidations to access the desired amino substituted thioamides is of considerable interest. Furthermore, it is also highly desirable to develop environmentally benign chemical processes without requirement of any metal catalyst and solvent-free conditions.

Recently, organo catalysts have been employed in a variety of chemical transformations [26-30] and they dominate the natural world in triggering chemical reactions. Particularly, L-proline has received much attention due to its dual role as a ligand and catalyst [31-34]. In view of the above perceptions, the development of benign and metal-free transamidation procedures with high yield and selectivity is desirable. In continuation of our interest on the development of environment-friendly transamidation catalysts [35-39], we wish to report a general L-proline-catalysed transthioamidation of primary amides with amines under solvent-free conditions [40]. To the best of our knowledge very rare reports available for the efficient transthioamidations under neat conditions [41].

For the initial studies, we chose thioacetamide 1a and benzyl amine 2a as substrates to explore the transthioamidations using L-proline as catalyst (Figure 1). Initially, when 1a and 2a were reacted with 5 mol % of L-proline catalyst in water at 130°C in a sealed tube, the desired transthioamidations derivative 3a was isolated in 10% yield after 36 h (Figure 1, entry 1). In ethanol as solvent 19% of 3a was isolated (Figure 1, entry 2). Shifting to other organic solvents (toluene, DMF, DMSO, NMP and DMA), the yield of the product was varied between 42% and 85% (Figure 1, entries 3-7). To our delight, the reaction was also very facile under neat conditions at 130°C and gave 3a in 89% yield (Figure 1, entry 8). Further, no improvement in the yield was observed either

*Corresponding author: Subbarayappa Adimurthy, Academy of Scientific and Innovative Research, CSIR–Central Salt and Marine Chemicals Research Institute, GB Marg, Bhavnagar-364002, Gujarat, India, Tel: 6860-2567760; E-mail: adimurthy@csmcri.org

Received April 18, 2016; Accepted May 09, 2016; Published May 17, 2016

Citation: Rao SN, Mohan DC, Adimurthy S (2016) L-Proline: An Efficient and Selective Catalyst for Transamidation of Thioamides with Amines. J Biomol Res Ther 5: 140. doi:10.4172/2167-7956.1000140

Copyright: © 2016 Rao SN, et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.
by lowering the reaction temperature or by increasing the catalyst loading (Figure 1, entries 9–12). Under the same conditions, without catalyst only 43% of desired product 3a was isolated (Figure 1, entry 13). Increasing the reaction temperature yield was not improved; decomposition of the product was observed (Figure 1, entries 14 and 15). Transmethioamidation was not efficient with other amino acid catalysts tested (Figure 1, entries 16–19).

With the set of optimized reaction conditions in hand, we moved on to investigate the scope of this metal-free transmethioamidation. A series of amines were subjected to the transmethioamidation of thioacetamide under these conditions (Figure 2). The reaction was found to be very facile with both electron-rich and moderately electron-deficient amines and produced corresponding transmethioamidation products 3a–3f in moderate to good yield (46–89%). The transmethioamidation was also efficient with variety of amines (alpha methyl, secondary benzyl, cyclic secondary, cyclohexyl, aryl alkyl and long chain aliphatic amines) and provided the corresponding products 3g–3n in moderate to good yield (59%–86%). Similarly, transmethioamidation of 2-(pyridin-2-yl) ethan-1-amine also gave 85% yield of desired product (3o).

To show the synthetic utility of this method, a variety of thioamides and amines were subjected to these optimized conditions (Figure 3). As expected, the transmethioamidation of thiobenzamide with variety of amines [benzyl amines (electron-neutral, -rich, -deficient), alkyl aromatic, aliphatic, cyclic secondary amines and hetero amines] provided the corresponding products 5a–5m in moderate to good yields (33%–82%). Similarly, hetero amines like pyridin-2-ylmethanamine as well as hetero thioamide were also gave the corresponding thioamidation (33%-82%). Similarly, hetero amines like pyridin-2-yl ethan-1-amine also gave 85% yield of desired product (3o).

Conclusion

In summary, we have reported the synthesis of variety of thioamides using easily available L-proline catalysed transamidation of various thioamides with amines under neat conditions. With this method a variety of corresponding thioamides were obtained in good to excellent yields under solvent-free conditions.

Acknowledgement

CSIR-CSMCR Communication No. 201/2015. We thank the “Analytical Discipline and Centralized Instrumental Facilities” for providing instrumentation facilities, S.N.R and D.C.M. are also thankful to CSIR and UGC, New Delhi for their fellowships. We thank DST, Government of India (SR/S1/OC-13/2011) and CSIR (CSC-0123) for the financial support.
Citation: Rao SN, Mohan DC, Adimurthy S (2016) L-Proline: An Efficient and Selective Catalyst for Transamidation of Thioamides with Amines. J Biomol Res Ther 5: 140. doi:10.4172/2167-7956.1000140

References

1. Jagodzinski TS (2003) Thioamides as useful synthons in the synthesis of heterocycles. Chem Rev 103: 197-228.
2. Danilkina NA, Mikhailov LE, Ivin BA (2006) Condensation of thioamides with acetylene carboxlic acid derivatives. Russian J Org Chem 42: 831-838.
3. Banala S, Süssmuth RD (2010) Thioamides in nature: in search of secondary metabolites in anaerobic microorganisms. Chem Bio Chem 11: 1355-1337.
4. Fang FG, Maier ME, Danishefsky SJ, Schultz G (1990) New routes to functionalized benzazepine substructures: a novel transformation of an alpha-diketone thioamide induced by trimethyl phosphite. J Org Chem 55: 831-838.
5. Nicolaou KC, Ligos DE, Kim DW, Noronha RG, et al. (2006) Total synthesis and biological evaluation of halipeptins A and D and analogues. J Am Chem Soc 128: 4460-4470.
6. Nicolaou KC, Delhe DH, Chen DXY (2008) Total syntheses of amythiamicins A, B and C. Chem Commun 2632-2634.
7. Padwa A, Beatt LS, Heidelbaugh TM, Liu B, Sheehan SM (2000) A one-pot bicycloanulation mechanism for the synthesis of tetrahydroisoquinoline systems. J Org Chem 65: 2684-2695.
8. Hu WP, Chen YK, Liao CC, Yu HS, Tsai YM, et al. (2010) Synthesis, and biological evaluation of 2-(4-aminoaryl)benzothiazole derivatives as photosensitizing agents. Bioorg Med Chem 18: 6197-6207.
9. Shi DF, Bradshaw TD, Wright S, McCull CJ, Leliiveld P, et al. (1996) Antitumor benzothiazoles 3. Synthesis of 2-(4-aminoaryl) benzothiazole and evaluation of their activities against breast cancer cell lines in vitro and in vivo. J Med Chem 39: 3375-3384.
10. Stevens MFG, McCull CJ, Leliiveld P, Alexander P, Richter JA, et al. (1994) Structural Studies on Bioactive Compounds. Synthesis of Polyhydroxylated 2-Phenylbenzothiazole and a Comparison of Their Cytotoxicities and Pharmacological Properties with Gentisine and Quercetin. J Med Chem 37: 1689-1695.
11. Fajardo TT, Guinto RS, Cellona RV, Abalos RM, Dela EC, et al. (2006) A clinical trial of ethionamide and prothionamide for treatment of lepromatous leprosy. Am J Trop Med Hyg 74: 457-461.
12. Akjo DM, Nasso PS, Hadley WK (1987) Therapeutic implications of inhibition versus killing of Mycobacterium avium complex by antimonials. Antimicrob Agents Chemother 31: 11-12.
13. Hwang J, Choi MG, Eor S, Chang SK (2012) Fluorescence signaling of Zr and C. Chem Commun 2632-2634.
14. Cava MP, Levinson MI (1975) Thionation reactions of lawesson’s reagents. Tetrahedron 41: 5061.
15. Zacharie B, Gilles Sauvé G, Penney C (1993) Enzymatic acylation and alkoxycarbonylation of α-, ω-, ω-hydroxy- and arboxy-nucleosides. Tetrahedron 49: 10089-10098.
16. Curphey T (2002) Thionation with the reagent of phosphorus pentasulfide and hexamethyldisiloxane. J Org Chem 67: 6461-6473.
17. Bergman J, Pettersson B, Hasimbegovic V, Svensson PH (2011) Thionations using a PS10-pyridine complex in solvents such as acetonitrile and dimethyl sulfones. J Org Chem 76: 1546-1553.
18. Shibahara F, Sugiiura R, Murai T (2009) Direct thionation and selenation of amides using elemental sulfur and selenium and hydrochlorosilanes in the presence of amine. Org Lett 11: 3064-3067.
19. Sztostak M, Aubé J (2009) Synthesis and rearrangement of a bridged thioamide. Chem Commun pp: 7122-7124.
20. Zbryuev OI, Sitasni N, Kappe CO (2003) Preparation of thioamide building blocks via microwave-promoted three-component kidnier reactions. J Comb Chem 5: 145-147.
21. Okamoto K, Yamamoto T, Kanbara T (2007) Efficient synthesis of thiobenzamides by Willgerodt-Kindler reaction with base catalysts. Synlett 2687-2690.
22. Nguyen TB, Ermolенко LAMA (2012) Efficient and selective multicomponent oxidative coupling of two different aliphatic primary amines into thioamides by elemental sulphur. Org Lett 14: 4274-4277.
23. Sun Y, Jiang H, Wu W, Zeng W, Li J (2014) Synthesis of thioamides via one-pot A3-coupling of alkyl bromides, amines, and sodium sulphide. Org Biomol Chem 12: 700-707.
24. Ji-Wei W, Ya-Dong W, Jian-Jun D, Hua-Jian X (2014) Benzoic acid-catalyzed transamination reactions of carboxamides, phthalimide, urea and thioamide with amines. Adv Synth Catal 356: 2429-2436.
25. Ojeda P, Gamba S (2015) Thioamide catalysis of transamination and transpeptidation reactions by StCl2. Tetrahedron Letters 56: 4308-4311.
26. Dalko PL, Moisan L (2004) In the golden age of organocatalysis. Angew Chem Int Ed 43: 5138-5175.
27. MacMillian DWC (2008) The advent and development of organocatalysis. Nature 455: 304-308.
28. Koalski H, Ikishima H, kuyama A (2008) Organocatalytic asymmetric synthesis using proline and related molecules. Part 2. Heterocycles 75: 757-797.
29. Karstelien E, Jorgensen KA (2009) Organocatalysis—after the gold rush. Chem Soc Rev 38: 2178-2189.
30. Chai Z, Zhao G (2012) Efficient organocatalysts derived from simple chiral acrylic amino acids in asymmetric catalysis. Catal Sci Technol 2: 29-41.
31. Couthard GL, Berg W, Aggarwal VK (2012) Stereocatalyzed organocatalytic synthesis of prostaglandin PFG2 in seven steps. Nature 489: 278-281.
32. Tanimori T, Ueno M, Takeda K, Kirihata M, Tanimori S (2012) Proline catalyzes direct C-H aminations of unactivated amines. J Org Chem 77: 7844-7847.
33. Tanimori S, Kobayashi Y, Iesaki Y, Ozaki Y, Kirihata M (2012) Copper-catalyzed synthesis of substituted indazoles from 2-chloroarenes at low catalyst-loading. Org Biomol Chem 10: 1381-1387.
34. Nezhad AK, Saniharghi S, Shahidzadeh ES, Panahi F (2012) L-Proline-promoted three-component reaction of anilines, aldehydes and barbituric acids/malononitrile: regioselective synthesis of 5-arylpiperidino [4,5-b] quinoline-diones and 2-amino-4-arylnitrones-3-carbonitriles in water. Green Chem 14: 2876-2884.
35. Rao SN, Mohan DC, Adimurthy S (2013) L-Proline: an efficient catalyst for transamination of carboxamides with amines. Org Lett 15: 1491-1493.
36. Rao SN, Mohan DC, Adimurthy S (2014) Chitosan: an efficient recyclable catalyst for transamination of carboxamides with amines under neat conditions. Green Chem 16: 4122-4126.
37. Rao SN, Mohan DC, Adimurthy S (2015) H-b-zeolite catalyzed transamination of carboxamides, phthalimide, formamides and thioamides with amines under neat conditions. RSC Adv 5: 95313-95317.
38. Mohan DC, Rao SN, Ravi C, Adimurthy S (2014) Copper(I) iodide Catalyzed Aerobic Oxidative C-N and C-S bond formations through C-H Activation: Synthesis of Functionalized Imidazo-[1,2-a]pyridines. Asian J Org Chem 3: 609-614.
39. Ravi C, Mohan DC, Adimurthy S (2014) N-Chlorosuccinimide-promoted regioselective sulfenylation of imidazoheterocycles at room temperature. Org Lett 16: 2978-2981.
40. Fioravanti S, Parise L, Pelagalli A, Pellecanni L, Trilli L (2015) Synthesis of N-benzoylthioacetamide (3a): In a sealed tube, 2.0 mmol of 1a, 2.0 mmol of 2a and L-proline (11.5 mg) were stirred at indicated temperature for indicated reaction time (See Schemes 1-2 and Table 1). After being cooled to room temperature, the reaction mixture was extracted with DCM (3 X 20 mL). After removal of solvent, the crude reaction mixture left out was purified by recrystallization or silica gel (200-400 mesh) column chromatographic separation (resolved in dichloromethane, eluted with dichloromethane and ethyl acetate). RSC Adv 5: 29312-29318.
41. Schlatter MJ (1942) The reaction between thioamides and primary amines. J Am Chem Soc 64: 2722-2723.

Scheme 1: Plausible reaction mechanisms.