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Practical chemoenzymatic synthesis of a 3-pyridylethanolamino β_3 adrenergic receptor agonist

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Abstract

A chemoenzymatic synthesis of β_3 agonist 1 suitable for large scale preparation is described. The key chiral 3pyridylethanolamine intermediate (*R*)-7 was prepared via an improved Neber rearrangement and a yeast-mediated asymmetric reduction. The tetrazolone fragment of the molecule was constructed via a dipolar cycloaddition between 1-(cyclopentyl)-3-propylazide and *p*-chlorosulfonyl phenylisocyanate. Sulfonamide coupling of these two intermediates under Shotten-Baumann conditions, followed by a borane reduction of the amide afforded 1 in 20-32% overall yield from 3-acetylpyridine. © 1999 Elsevier Science Ltd. All rights reserved.

Obesity affects approximately 30% of the adult population, and is closely associated with the development of type II diabetes, coronary artery diseases, and hypertension. The morbidities associated with these diseases are sometimes reversed by weight loss.¹ Compound 1 is proposed to act by a novel mechanism elevating metabolic rate through thermogenesis resulting from the stimulation of β_3 adrenergic receptors (AR) in brown adipose tissue.² The elevation of metabolic rate, in the absence of increased food intake, will lead to weight loss over time. β_3 AR agonists have also demonstrated a direct improvement on glucose tolerance and therefore may be very important in the treatment of type II diabetes.¹ In order to carry out further studies with this compound, we required a practical chiral synthesis. This paper describes the process used to provide kilogram quantities of compound 1, a potent, selective β_3 AR agonist.³

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Compound 1 consists of a chiral *N*-alkylated 3-pyridylethanolamine fragment and a tetrazolone sulfonyl fragment connected through a sulfonamide bond. Previous synthesis of the pyridyl ethanolamine portion involved a (-)-DIP-Cl asymmetric reduction of 3-pyridyl chloromethyl ketone, followed by epoxide formation of the resulting chlorohydrin, and then opening of the epoxide with the appropriate amine.² For our synthesis, we took a different approach where the amine moiety was incorporated in the form of amide early in the synthesis prior to introduction of the chiral alcohol. The synthesis of 7 is outlined in Scheme 1.



Scheme 1. a. 1. NH₂OH·HCl; 2. TsCl, pyr; b. 1. EtOK, EtOH; 2. HCl, MTBE; c. EDC, THF, H₂O, pH 5.5, *p*-nitrophenyl acetic acid; d. aq. HCl; e. yeast-mediated bioreduction; f. H2, Pd/C, MeOH, NH₄OH

We have improved significantly upon an organic synthesis procedure⁴ for the preparation of 3-pyridylaminomethyl ketal **3** via a Neber rearrangement reaction, and also noted some safety issues.⁵ Compound **3** was prepared in 76% overall yield from 3-acetylpyridine in three steps via tosyl oxime **2**. The oxime formation and the tosylation steps⁵ (93% overall yield) were carried out in a one-pot fashion using pyridine as the solvent. Under such conditions, the *E*-isomer of oxime predominates (>97:3). Water was removed by azeotropic distillation prior to the tosylation. Tosyl oxime **2** was crystallized from the reaction mixture by addition of water, and the wet cake was used directly in the Neber rearrangement by adding EtOK to a mixture of **2** in EtOH at 10–30°C. After removal of the TsOK salt,⁶ the filtrate was treated with gasesous HCl (final pH 1–2) to precipitate the amino ketal di-HCl salt **3** (82%). A carbodiimide-mediated coupling of **3** with *p*-nitrophenylacetic acid in THF/H₂O followed by a selective hydrolysis of the resulting amide-ketal **4** with concomitant removal of ethanol afforded pyridyl ketone **5** (93% for the two steps).¹²

With ketone 5 in hand, we screened methods for the key asymmetric reduction of the ketone to the chiral alcohol. Unfortunately, preliminary results from most known methods gave poor enantioselectivity and in some cases reduction of pyridine ring was observed (e.g. asymmetric hydrogenation with (S)-tol-BiNAP--RuCl₂--Et₃N complex afforded the reduced pyridine vinyloguous amide as the major product, with 6 as the minor product in only 6% ee). Alternatively, initial results using Baker's yeast (Saccharomyces cerevisiae) were encouraging,⁷ which provided product in 70-80% ee. Subsequently, a microbial screen identified the yeast Candida sorbophila as a suitable biocatalyst,⁸ which furnished the

chiral alcohol 6^{12} in ~98% ee and 70–90% conversion after incubation for 2 days at 28°C. The material was routely isolated in ~75% yield and >99.5% ee.⁹ The absolute configuration of the secondary alcohol produced in the yeast-mediated reduction of ketone **5** was determined to be the desired *R*-enantiomer by NMR analysis of the (*S*)-methoxyphenylacetate esters based on Trost's method¹⁰ and by comparision with materials derived from Shih's synthesis using (–)-DIP-Cl.² As shown below, the chemical shifts of the pyridyl protons of the *R*-isomer was shielded by the mandelate phenyl ring by 0.11–0.25 ppm relative to the *S*-isomer (at +20°C).

The optical purities of the crude **6** in the fermentation broth and the isolated **6** were determined to be >97.5% ee and >99.5% ee, respectively, by chiral HPLC using Chiracel OD-H (20/80 IPA/hexanes, 0.8 mL/min, 254 nm; or SFC, 300 bar CO₂, 16% MeOH (20 mM *i*PrNH₂), 1.0 mL/min, 35°C).



Nitro compound **6** was hydrogenated (20 psi H₂) over 5% Pd/C (4 wt%) in methanol with 1.7 equiv. NH₄OH, conditions which minimized impurities derived from the intermediary nitroso compound. The reaction was run at 40°C for \sim 3 h until the uptake of H₂ had stopped. Aniline **7** was crystallized in high yield (96–97%) and purity (>99 A%).¹²

The southern half of the molecule was constructed as shown in Scheme 2. The cycloaddition between azide 13 and isocyanate 15 took place at 125°C to provide the tetrazolone 8 in 84% yield.¹¹



Scheme 2.

Subsequent coupling of aniline 7 and sulfonyl chloride 8 under Schotten–Baumann conditions afforded the penultimate amide 9 in 94% yield (Scheme 3). Selective reduction of the amide with $BH_3 \cdot SMe_2$ (7 equiv.) at +20°C followed by 1 M H₂SO₄ work up at +50°C to break up the borane complexes afforded the crude 1 free base in 82% yield after neutralization. This material was purified to >99% pure by crystallization of the corresponding phosphate salt (93%) followed by free basing (98%), since the di-HCl salt gave poor rejection of the impurities. Finally, the di-HCl salt was prepared from ethanolic HCl in 92% yield.¹²

In summary, we have developed a practical synthesis of β_3 AR agonists 1 and have demonstrated it on kilogram scale. The key chiral pyridyl alcohol intermediate (*R*)-7 was prepared via an improved Neber rearrangement and a yeast-mediated asymmetric reduction. The key dipolar cycloaddition between azide 15 and isocyanate 17 provided an efficient one-step synthesis of the tetrazolone sulfonyl chloride.



Scheme 3.

Coupling of the aniline and the sulfonyl chloride followed by borane reduction of the amide completed the synthesis of 1 in 20–32% overall yield from 3-acetylpyridine.

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- All the new compounds were well characterized by ¹H NMR, ¹³C NMR and elemental analysis. Data for compounds **3**, **5**, **7**, **8**, and **1** are listed below. For compounds **2**, **4**, **6**, and **9** data are readily available upon request. Amino ketal **3**: ¹H NMR (CD₃OD) δ 9.07 (d, J=1.9 Hz, 1H), 8.99 (d, J=6.0 Hz, 1H), 8.83 (dt, J=8.3, 1.6 Hz, 1H), 8.26 (dd, J=8.3, 6.0, 1H), 3.61 (m, 2H), 3.59 (brs, 2H), 3.46 (m, 2H), 1.30 (t, J=7 Hz, 6H); ¹³C NMR (CD₃OD) δ 147.1, 143.8, 142.5, 139.9, 129.3, 99.9, 59.2, 44.5, 15.0. Anal. calcd for C₁₁H₂₀Cl₂N₂O₂·3/4 KCl: C, 38.96; H, 5.94; N, 8.26. Found: C, 38.81; H, 5.83; N, 8.00. Keto-amide **5**: ¹H NMR (CDCl₃) δ 9.17 (d, J=1.7 Hz, 1H), 8.84 (dd, J=4.8, 1.7 Hz, 1H), 8.22 (d, J=8.7 Hz, 2H), 8.23 (m, 1H), 7.52 (d, J=8.7 Hz, 2H), 7.47 (m, 1H), 6.60 (brs, NH), 4.78 (d, J=4.4 Hz, 2H), 3.78 (s, 2H); ¹³C NMR (CDCl₃) δ 193.1, 169.3, 154.7, 149.3, 147.3, 141.8, 135.2, 130.3, 129.7, 124.0, 123.9, 46.7, 43.0. Anal. calcd for C₁₅H₁₃N₃O₄: C, 60.20; H, 4.38; N, 14.04. Found: C, 59.82; H, 4.24; N, 13.83. Aniline-alcohol 7: ¹H NMR (CD₃OD) δ 8.49 (d, J=2.0 Hz, 60.20; H, 4.38; N, 14.04. Found: C, 59.82; H, 4.24; N, 13.83. Aniline-alcohol 7: ¹H NMR (CD₃OD) δ 8.49 (d, J=2.0 Hz, 60.20; H, 4.38; N, 14.04. Found: C, 59.82; H, 4.24; N, 13.83. Aniline-alcohol 7: ¹H NMR (CD₃OD) δ 8.49 (d, J=2.0 Hz, 60.20; H, 4.38; N, 14.04. Found: C, 59.82; H, 4.24; N, 13.83. Aniline-alcohol 7: ¹H NMR (CD₃OD) δ 8.49 (d, J=2.0 Hz, 60.20; Hz, 70.20; Hz,

1H), 8.40 (dd, J=4.9, 1.6 Hz, 1H), 7.72 (dt, J=8.0, 1.6 Hz, 1H), 7.34 (dd, J=7.8, 5.0 Hz, 1H), 6.94 (d, J=8.4 Hz, 2H), 6.66 (d, J=8.4 Hz, 2H), 4.78 (t, J=6.1 Hz, 1H), 3.42 (d, J=6.1 Hz, 2H), 3.31 (s, 2H); ¹³C NMR (CD₃OD) δ 175.2, 149.1, 148.3, 147.7, 140.2, 136.3, 130.8, 125.9, 125.1, 116.8, 71.1, 47.5, 43.1. Anal. calcd for C₁₅H₁₇N₃O₂: C, 66.40; H, 6.32; N, 15.49. Found: C, 66.07; H, 6.35; N, 15.29. Tetrazole sulfonyl chloride **8**: ¹H NMR (CDCl₃) δ 8.35 (d, J=9.1 Hz, 2H), 8.16 (d, J=9.1 Hz, 2H), 4.03 (t, J=7.2 Hz, 2H), 1.90 (m, 2H), 1.85–1.70 (m, 3H), 1.65–1.45 (m, 4H), 1.40 (m, 2H), 1.08 (m, 2H); ¹³C NMR (CDCl₃) δ 148.7, 142.2, 139.9, 128.7, 118.8, 45.6, 39.5, 32.7, 32.6, 27.6, 25.1. Anal. calcd for C₁₅H₁₉ClN₄O₃S: C, 48.58; H, 5.16; Cl, 9.56; N, 15.11. Found: C, 48.87; H, 5.21; Cl, 9.29; N, 14.90. 1-2HCl: ¹H NMR (CD₃OD) δ 9.01 (d, J=1.2 Hz, 1H), 8.86 (d, J=5.6 Hz, 1H), 8.76 (dt, J=8.1, 1.5 Hz, 1H), 8.14 (dd, J=8.1, 5.6 Hz, 1H), 8.06 (d, J=8.9 Hz, 2H), 7.88 (d, J=8.9 Hz, 2H), 7.19 (d, J=8.6 Hz, 2H), 7.10 (d, J=8.6 Hz, 2H), 5.37 (dd, J=9.8, 3.1 Hz, 1H), 4.00 (t, J=7.1 Hz, 2H), 3.49 (dd, J=12.8, 3.3 Hz, 1H), 3.35–3.20 (m, 3H), 3.00 (m, 2H), 1.92–1.70 (m, 5H), 1.70–1.45 (m, 4H), 1.45–1.30 (m, 2H), 1.10 (m, 2H); ¹³C NMR (CD₃OD) δ 150.1, 145.9, 143.3, 142.5, 141.1, 139.7, 139.4, 137.8, 134.4, 130.7, 129.7, 128.7, 122.8, 120.0, 66.9, 53.7, 50.0, 46.4, 40.9, 33.9, 33.6, 32.4, 28.6, 26.1. Anal. calcd for C₃₀H₃₉C₁₂N₇O₄S: C, 54.21; H, 5.91; Cl, 10.67; N, 14.75; S, 4.82. Found: C, 54.35; H, 5.90; Cl, 10.78; N, 14.64; S, 4.87.