Synthesis and Characterization of Bromoclenbuterol

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Abstract
Clenbuterol is a sympathomimetic amine used for breathing disorders as a decongestant and bronchodilator. During the process development of clenbuterol, the process related impurity bromoclenbuterol was identified as a critical impurity along with the final API. The present work describes the synthesis and characterization of this bromoclenbuterol.

Keywords: B ICH; Breathing disorders; Impurity; Bromoclenbuterol

Introduction
Clenbuterol, it is most commonly available as the hydrochloride salt, clenbuterol hydrochloride. Clenbuterol, marketed as Dilaterol, Spiropent, Ventipulmin, and also generically as clenbuterol, is a sympathomimetic amine used for breathing disorders as a decongestant and bronchodilator. People with chronic breathing disorders such as asthma use this as a bronchodilator to make breathing easier. Clenbuterol is a β2 agonist with some structural and pharmacological similarities to epinephrine and salbutamol, but its effects are more potent and longer-lasting as a stimulant and thermogenic drug. It causes an increase in aerobic capacity, central nervous system stimulation, blood pressure, and oxygen transportation. It increases the rate at which body fat is metabolized while increasing the body’s BMR. It is commonly used for smooth muscle-relaxant properties as a bronchodilator and tocolytic. Clenbuterol is also prescribed for treatment of horses, but equine use is usually the liquid form (Figure 1).

Clenbuterol Hydrochloride was first synthesized at Thomae; a Boehringer Ingelheim research facility in Biberach, Germany, in 1967. The synthesis of Clenbuterol Hydrochloride was patented in the United States in 1970. After comprehensive clinical trials, Clenbuterol Hydrochloride was approved for the treatment of reversible airway obstruction in Germany in 1976 and later as a veterinary pharmaceutical for the treatment of bronchiolytic disorders in Germany in 1980. Boehringer Ingelheim markets Clenbuterol Hydrochloride as Spirospent for Human Pharmaceuticals and as Ventipulmin for Veterinary Pharmaceuticals. Clenbuterol Hydrochloride is not approved by the Federal Drug Administration for human use in the United States.

Impurity is defined as any substance coexisting with the original drug, such as starting material or intermediates or those formed, due to any side reactions. The presence of these unwanted chemicals, even in small amounts, may influence the efficacy and safety of the pharmaceutical products. A general scheme is set for the estimation of the impurities in bulk drug substances by the rational use of chromatographic, spectroscopic and analytical techniques. The process-related impurities in an active pharmaceutical ingredient (API) can have a significant impact on the quality and safety of the drug products. It is quite important for “regulatory” aspect of drug approval to provide limitation of “related impurities.” Therefore, it is necessary to study the impurity profile of any API and control it during the manufacturing of a drug product. As per the ICH guidelines, impurities which are forming at a level of ≥ 0.10% with respect to the API should be identified, synthesized, and characterized thoroughly [1,2].

During the process development of clenbuterol hydrochloride in RA Chem Pharma Ltd [3], the process related impurities as per pharmacopoeia (British pharmacopoeia and European pharmacopoeia) are identified and they are listed as 4-amino-3,5-dichloro benzaldehyde (Impurity-A), 1-(4-amino-3,5-dichlorophenyl)-2-[(1,1-dimethylethyl)amino]ethanone (Impurity-B), 1-(4-amino-3,5-dichlorophenyl)ethanone (Impurity-C), 1-(4-aminophenyl)ethanone (Impurity-D), 1-(4-amino-3,5-dichlorophenyl)-2-bromo ethanone (Impurity-E) and (1RS)-1-(4-amino-3-bromo-5-chloro phenyl)-2-(1,1-dimethylethyl)amino ethanol or bromoclenbuterol (Impurity-F).

The purpose of this work is for determining the impurities of clenbuterol to ensure the quality, efficacy and safety of the active ingredient in final pharmaceutical formulation. Among the six process related impurities, the bromoclenbuterol impurity was identified as a critical impurity and it was formed in the synthesis of clenbuterol hydrochloride API at the time of chlorination of 4-amino acetophenone (Table 1).

The present article describes a simple and facile synthesis for bromoclenbuterol impurity. This may serve as a standard for impurity profiling in drug development and analytical method validations.

Experimental
The chemicals and solvents were purchased from commercial suppliers and they were used without purification prior to use. 1H NMR spectra were recorded on 400 MHz and 13C NMR spectra were recorded on 100 MHz FT-NMR spectrometers. All chemical shifts

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are given as δ value with reference to Tetra methyl silane (TMS) as an internal standard.

1-(4-Amino-3-chlorophenyl)ethanone (7)

To a stirred solution of 1N HCl (1500 ml) was added 4-amino acetophenone (1) (200 gm, 1.48 mole) and N-Chloro succinimide (50 gm, 0.37 mole) at room temperature, and stirring continued for 3 hrs at 25-30°C. After maintenance undissolved material was filtered from the reaction mixture, total filtrate was taken and extracted with ethyl acetate, dried over sodium sulphate and evaporated under vacuum to get crude. Crude material was dissolve in ethyl acetate, titrated with ethyl acetate, and this acidic titration operation was repeated 2 times to get mono chloro compound as solid material, this solid material was neutralized with sodium carbonate solution in aqueous condition and further purified by using recrystallization technique in ethyl acetate to get 68 gm (yield-27%) 3-chloro-4-amino acetophenone (7) (mono chloro compound), as light brown colored solid with 98.66% HPLC purity (124 gm of unreacted 4-aminocacetophenone obtained from aqueous layer).

1-(4-Amino-3-bromo-5-chlorophenyl)-2-bromoethanone (8)

To a stirred solution of 3-chloro-4-amino acetophenone (7) (14 gm, 0.082 mole) in chloroform (140 ml) was added bromine (26.24 gm, 0.164 mole) solution slowly at 25-30°C and stirring continued for 6 hrs at same temperature. After completion of the reaction, methanol was added to the reaction mixture and continued the stirring for 30 min at RT. Undissolved material was filtered, the filtrate was distilled up to 50%, remaining mass was cooled to 0-5°C and filtered to give 15 gm (yield-55%) of 1-(4-amino-3-chloro-5-bromo-phenyl)-2-bromoethanone (8) as light brown color solid with 95.15% HPLC purity.

2-(Tert-butylamino)-1-(4-amino-3-bromo-5-chlorophenyl)ethanone (9)

To a stirred solution of 1-(4-amino-3-chloro-5-bromo-phenyl)-2-bromoethanone (8) (8 gm, 0.024 mole) in chloroform (50 ml) was added catalytic amount of potassium iodide (0.1 gm, 0.0006 mole) and tertiary butyl amine (5.2 gm, 0.072 mole) at 0-5°C and stirring was continued for 24 hrs at 0-5°C. After completion of the reaction, undissolved salts were filtered, the filtrate was distilled under vacuum to get crude solid material, which was triturated with hexane to give 6 gm (yield-76%) of 1-(4-amino-3-chloro-5-bromo-phenyl)-2-[(1,1-dimethylethyl)amino]ethanone (9) as light pale yellow color solid.

(S)-2-[(Tert-butylamino)-1-(4-amino-3-bromo-5-chlorophenyl)ethanol (10)

To a stirred solution of 1-(4-amino-3-chloro-5-bromo-phenyl)-2-[(1,1-dimethylethyl)amino]ethanone (9) (6 gm, 0.018 mole) in methanol (25 ml) was added sodium borohydride (0.7 gm, 0.018 mole) at 0-5°C. After addition, reaction mixture was slowly allowed to come to room temperature and stirred for 10 hrs at 25-30°C. On completion, reaction mixture was poured in to chilled water, obtained precipitate was filtered, dried and recrystallized in methanol to give 5 gm (yield-82%) of 1RS-1-(4-amino-3-bromo-5-chlorophenyl) -2-[(1,1-dimethyl ethyl)amino]ethanol (or) Bromo clenbuterol (10) as off-white solid. HPLC purity-98.80%, 1H NMR (CDCl₃): δ 7.35 (d, 1H, J=1.2 Hz), 7.23 (d, 1H, J=1.6 Hz), 4.45 (br s, 2H), 4.42 (dd, 1H, J=9.2, 3.6 Hz), 2.84 (dd, 1H, J=11.6, 3.6 Hz), 2.50 (dd, 1H, J=12.0, 9.2 Hz), 1.10 (s, 9H), 13C NMR (CDCl₃): 140.12, 133.93, 128.46, 126.05, 119.16, 109.08, 70.94, 50.33, 50.05, 29.15. IR (KBr, Cm−1): 3465.95, 3320.19, 2965.04, 1623.40, 1483.88, 1219.17, 758.77, 630.41. Mass: (m/z)-323.01 (M+• peak).

Results and Discussion

As per the available literature [4-7], clenbuterol hydrochloride was synthesized from 4-amino acetophenone (Scheme 1). Initially 4-amino acetophenone (1) was reacted with chlorine to afford 4-amino-3,5-dichloro acetophenone (2) which was further reacted bromine to give...
1-(4-amino-3,5-dichlorophenyl)-2-bromoethanone (3). The obtained bromo compound was reacted tertiary butyl amine to afford 2-(tert-butylamino)-1-(4-amino-3,5-dichlorophenyl)ethanone (4), which was further reduced with sodium borohydride to give clenbuterol base (5) and converted to hydrochloride salt by using alcoholic HCl to get clenbuterol hydrochloride (6).

In order to find the control and prevention of process related impurities of clenbuterol, the preliminary efforts were mainly focused on the manufacturing process. Among the six process related impurities, the bromoclenbuterol impurity was identified as a critical impurity because it cannot be eliminated from the API molecule, after formation in the reaction. Bromoclenbuterol and clenbuterol possessed same properties like solubility and crystallization nature. We turned our attention to identify the root cause for the formation of bromoclenbuterol in clenbuterol synthetic process. After thorough investigation, it was identified as mono chloro intermediate is the main root cause for the bromoclenbuterol formation.

In the synthesis of clenbuterol hydrochloride, first step was a double chlorination of 4-aminoacetophenone (1) through an electrophillic aromatic substitution reaction to yield 4-amino-3,5-dichloroacetophenone (2). Due to the ortho/para directing, amino group and the meta directing, electron withdrawing, acetyl group, chlorination of 4-aminoacetophenone occurs primarily at the 3 and 5 positions over the 2 and 6 positions. Therefore, under chlorination would produce only the mono chlorinated impurity, 4-amino-3-chloroacetophenone. Under these conditions, over chlorination does not result in the addition of chlorine to the 2 and 6 position because the amino and acetyl groups do not direct that addition. Even though chlorides are ortho/para directing and direct to the 2 and 6 position, chlorides are also deactivating. After close observation on this chlorination reaction, it was noted that the formed mono chlorinated impurity (Scheme 2) (4-amino-3-chloro acetophenone) caused the formation of process related impurity (bromoclenbuterol) in clenbuerol synthesis.

As per the above said observations on chlorination reaction, we concluded that the control of monochloro impurity should be itself in the initial stage i.e., chlorination reaction, by using different crystallization techniques. In the same way, the focus turned on the synthesis of bromoclenbuterol from mono chlorinated impurity by using Scheme 3 and this impurity should be necessary in the method development and validation of clenbuterol hydrochloride API molecule.

As per the above said observations on chlorination reaction, 4-Amino acetophenone (1) was reacted with N-Chlorosuccinimide in 1N HCl to afford 4-amino-3-chloro acetophenone (7), which was...
reacted with bromine to give 1-(4-amino-3-bromo-5-chlorophenyl)-2-bromoethanone (8). The obtained bromo compound was reacted with tertiary butyl amine to afford 2-(tert-butylamino)-1-(4-amino-3-bromo-5-chlorophenyl)ethanone (9), which was reduced with sodium borohydride in methanol to give bromoclenbuterol compound (10). The synthesized bromoclenbuterol structure was confirmed by $^1$H NMR, $^{13}$C NMR, IR and mass spectra.

**Conclusion**

The known impurity, bromoclenbuterol (Impurity-F in clenbuterol as per BP and EP) was synthesized and confirmed by the characterization tools such as HPLC, $^1$H NMR, $^{13}$C NMR, IR and MASS.

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