A Scalable, Chromatography-Free Synthesis of Benzotetramisole

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Abstract The scalable, chromatography-free synthesis of the chiral isothiourea benzotetramisole (BTM) in two steps from commercially available materials is presented. A detailed procedure for the synthesis of both enantiomers and the racemate on ca. 10 gram scale is disclosed.

Key words isothiourea, Lewis base catalysis, asymmetric catalysis, kinetic resolution, rearrangement, benzotetramisole

Introduction

Isothioureas are an important class of Lewis base that are becoming increasingly used within a range of organocatalytic procedures.1 This molecular class came to prominence in organocatalysis with the seminal report by Birman and Li in 20062 that demonstrated their power as acyl transfer catalysts in the kinetic resolution of secondary alcohols. The most potent and selective catalyst in this study was benzotetramisole (BTM), the benzannulated derivative of the antihelminthic pharmaceutical tetramisole3 that itself is a competent catalyst for this transformation. Based upon the isothiourea catalyst motif, further derivatisations of the heterocyclic core, including changes to ring size and substitution pattern, have led to a series of powerful dihydropyrimido[2,1-b]benzothiazole-based catalysts that have been applied to a range of transformations (Figure 1).4

Figure 1 A selection of isothiourea catalysts

The power of BTM (4) as a highly enantioselective acyl transfer catalyst has been further demonstrated by Birman,5 Shiina,6 and others7 in a range of kinetic resolution processes. Similarly, BTM has found use in an asymmetric Steglich rearrangement8 and the kinetic resolution of secondary alcohols by silylation.9 The reports by Romo and co-workers on the nucleophile-catalysed aldol-lactonisation

Scheme 1 A practical, multigram synthesis of benzotetramisole (BTM; 4)

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reaction using ammonium enolates, generated initially from cinchona alkaloids and later using isothioureas, stimulated our group to further demonstrate that isothioureas serve as excellent Lewis base organocatalysts for these processes. BTM (4) in particular has enabled the catalytic enantioselective synthesis of dihydropyridones and as well as promoting the asymmetric [2,3]-rearrangement of allylic ammonium ylides. Romo and co-workers have also reported the use of BTM (4) in the asymmetric synthesis of bicyclic lactones.

While tetrámisole is commercially available at reasonable cost and a scalable route to HBTM-2.1 has been reported, in our hands, Birman’s original procedure (detailed in Scheme 2) was suitable for the synthesis of (R)-BTM on ca. 1 g scale. Although this sequence was reproducible, in practice we found this sequence to be limited by the use of a sealed tube in the first step, chromatographic purifications to obtain both (R)-3 and noncrystalline (R)-4, and multiple recrystallisations to obtain pure material.

Further syntheses of BTM have been reported based upon this original procedure. Incremental improvements have been made by Chen and co-workers, finding firstly that intermediate 3 can be purified by recrystallisation from chloroform and secondly that the two reactions could be performed sequentially in ‘one-pot’ without isolation of 3. However, chromatography and recrystallisation were still required to furnish pure BTM and the final yield of crystalline material was not disclosed. Recently, Okamoto and co-workers have also reported the use of BTM (4) in the asymmetric synthesis of BTM (4).

Although (R)- and (S)-4 are now commercially available (ca. £ 90/1 g), the need for multi-gram quantities of racemic and enantioenriched BTM (4) to support our investigations into its use in catalysis motivated the development of a scalable, operationally simple synthesis. Herein we report the multi-gram synthesis of (R)-, (S)- and (±)-4 from commercially available 2-chlorobenzothiazole (1) and 2-phenylglycinol (2) without recourse to chromatographic purifications (Scheme 1).

**Scope and Limitations**

First, the neat reaction between 2-chlorobenzothiazole (1), 2-phenylglycinol (2) and ethyldiisopropylamine to form 3 was optimised. The use of a high-boiling co-solvent (chlorobenzene at reflux) whilst maintaining high concentrations of 1 (ca. 2.0 M) allowed the reaction to be performed in standard glassware under air without the need for sealed tubes or an inert atmosphere. However, extended reaction times (>48 h) at reflux (ca. 140 °C) were required for high conversions of 1 as measured by GC analysis. Switching to 1,2-dichlorobenzene and increasing the reflux temperature (ca. 195 °C) led consistently and reproducibly to >95% conversion of 1 after 24 hours (Scheme 3, a). Reducing the temperature, equivalents of ethyldiisopropylamine, or diluting the reaction further led to lower conversions and extended reaction times.

Chen and Nagano observed precipitation of product 3 in the separatory funnel during aqueous workup of the reaction. Seeking to exploit this to improve the workup procedure, the reaction mixture was diluted with water resulting in the formation of a thick paste. Upon addition of an organic solvent (e.g., hexanes or dichloromethane), large amounts of a fine solid precipitate of 3 was formed. This could be collected by filtration, followed by washing with further portions of organic solvent to leave crude 3 as a tan solid. In our hands, recrystallisation from hot chloroform as detailed by Chen returned the chloroform adduct of 3, from which residual chloroform could not be removed even after extended periods under reduced pressure. Multiple cycles of re-slurrying the solid with dichloromethane followed by evaporation of the solvent allowed the isolation of solid 3 that was pure by 1H and 13C{'H} NMR analysis (Scheme 3, b).

This material contained variable amounts of water as evidenced by droplets present during recrystallisation and slurring. This water was not purged by filtration and had a detrimental effect on the subsequent step. The azeotropic removal of water from 3 was achieved through heating a suspension of 3 at reflux in toluene using a Dean–Stark apparatus. However, 3 was observed to dissolve in toluene at reflux, so direct recrystallisation of crude 3 from toluene in a Dean–Stark apparatus was considered as a possible sim-
plification to access analytically pure, anhydrous \(3\). Using this process and starting from 10.00 grams of (R)-, (S)-, or (±)-2,\(^{18}\) this procedure gave analytically pure, fluffy white crystals of \(3\) consistently in 74–79% yield after one or two recrystallisations from hot toluene in a Dean–Stark apparatus (Scheme 3, c).\(^{19}\)

With large quantities of \(3\) easily accessible, the transformation of the alcohol functionality and cyclisation into BTM 4 was examined. Employing Birman’s conditions of methanesulfonyl chloride and triethylamine (Scheme 2) followed by aqueous workup yields a yellow gum which, although relatively pure by \(^1\)H NMR analysis, proved difficult to transform into high yields of crystalline BTM by trituration or recrystallisation without recourse to chromatography. This tentatively suggested the presence of polymeric impurities that were largely silent by \(^1\)H NMR analysis. Therefore, alternative activation methods for the alcohol cyclisation were examined.

Both tosyl chloride and thionyl chloride gave unsatisfactory results, with the former giving a complex mixture of products and the latter incomplete conversion of \(3\) (Scheme 4). Interestingly, the reaction of \(3\) under Mitsunobu conditions resulted in skeletal rearrangement to give isomeric isothiourea 5 as the major species. The structure was inferred by comparison of the crude \(^1\)H NMR spectra to that reported by Okamoto and co-workers.\(^4\) This unexpected reactivity was not pursued further, and the fidelity of stereochemical transfer was not determined.

As alternative activating agents were ultimately unsuccessful, efforts were focused upon understanding and improving the reaction of \(3\) with methanesulfonyl chloride. It was found that >1.25 equivalents of methanesulfonyl chlo-ride were required for complete consumption of \(3\), even with freshly distilled methanesulfonyl chloride under strictly anhydrous conditions. TLC and \(^1\)H NMR analysis of the reaction mixture indicated the formation of two products. Attempted chromatographic isolation of the two species failed to allow isolation of purported O-mesylate 6 owing to its instability on silica gel. However, a second N,O-bis-mesylate species 7 was isolated and characterised (Scheme 5). Treating \(3\) with >2 equivalents of methanesulfonyl chloride allowed 7 to be produced exclusively without any of the alternative isomeric N,O-bis-mesylate being observed.\(^{20}\) Interestingly, heating a dichloromethane solution of 7 at reflux in the presence of triethylamine and methanol overnight gave BTM 4 as the sole product in excellent yield after chromatography. This observation suggests that under Birman’s original conditions a mixture of mono- and bis-mesylate 6 and 7 forms with complete consumption of
methanesulfonyl chloride. Both components of this mixture cyclise upon heating with base to give 4 exclusively, with the sulfene eliminated from the cyclisation of bis-mesylate 7 quenched by methanol.\(^\text{21}\) Indeed, if a nucleophilic co-solvent is absent the reaction turns black and the isolable yield of BTM 4 is significantly reduced.

With this information in hand, optimised conditions for the cyclisation were established (Scheme 6). A suspension of 3 and excess triethylamine (4.0 equiv) in anhydrous dichloromethane (0.1 M) was cooled in an ice/water bath. A slight excess of methanesulfonyl chloride (1.3 equiv) was added, sufficient to completely consume 3. The resulting solution was treated with isopropanol,\(^\text{22}\) heated to reflux overnight, and checked for completion by \(^1\)H NMR analysis. These conditions consistently led to full conversion of 3 into crude 4. An improved workup procedure was sought to avoid chromatography, with the Brønsted basic nature of 4 considered an exploitable property through acid/base extractions. Hence, the reaction was quenched with aqueous 1 M sodium hydroxide to remove acidic impurities (MsOH, etc.) and the dichloromethane layer extracted multiple times with aqueous 1 M hydrochloric acid to leave an aqueous solution of BTM·HCl (\(\times\)times with aqueous 1 M hydrochloric acid to leave an aqueous layer. A solvent switch from dichloromethane to diethyl ether (in which BTM·HCl is only sparingly soluble) was performed after the aqueous 1 M sodium hydroxide wash and this greatly increased the mass recovery of crude 4 after the acid/base extractions. Final modifications to the workup included a toluene azeotrope during the solvent switch to remove the remaining triethylamine, and treatment of the final organic solution of 4 with charcoal to decolourise it. Finally, trituration of the crude with hot diethyl ether provided analytically pure BTM (4) in consistently high yield for both enantiomers and the racemate of 73–84% from 3 (Scheme 6).

In conclusion, the scalable, chromatography-free, and operationally simple synthesis of the isothiourea organocatalyst BTM has been demonstrated. The transformation of intermediate 3 with methanesulfonyl chloride has also been investigated, showing that both mono- and bis-mesylates 6 and 7 will cyclise to form BTM under the reaction conditions.

Reactions were performed in oven-dried glassware, using DrySyn\textsuperscript® blocks for heated reactions. Anhydrous CH\(_2\)Cl\(_2\) was obtained from an MBraun SPS-800 system. 2-Chlorobenzothiazole was fractionally distilled under reduced pressure prior to use. \(\text{i-Pr}_2\)NEt and Et\(_3\)N were stored over KOH pellets, and MsCl was distilled under reduced pressure from P\(_2\)O\(_5\) prior to use and stored in a fridge under N\(_2\).\(^\text{23}\) All other solvents and commercial reagents were used as received without further purification. Petroleum ether (PE) used refers to the fraction boiling in the 40–60 °C range. Analytical TLC was performed on precoated aluminum plates (Kieselgel 60 F\(_{254}\) silica). Plates were visualised under UV light (254 nm) or by staining with KMnO\(_4\) followed by heating. Flash column chromatography was performed on Kieselgel
A 250 mL round-bottomed flask containing a stirrer bar was charged with (R)-2-phenylglycinol ([R]-2; 10.00 g, 72.80 mmol, 1.05 equiv), o-dichlorobenzene (34.5 mL, 2.0 M), i-Pr2NEt (31.0 mL, 173.6 mmol, 2.5 equiv), and 2-chlorobenzothiazole (1; 8.60 mL, 69.43 mmol, 1.0 equiv. The resulting yellow suspension was stirred vigorously and heated to reflux at 195 °C (DrySyn® temperature), at which point the solid was suspended. The mixture was diluted with distilled H2O (100 mL) and toluene (75 mL) with vigorous stirring. A precipitate formed over the next 15 min and stirring was maintained for a further 1 h. The reaction mixture was filtered through a 1.0 L sintered glass filter funnel (porosity 3), and washed with further portions of toluene (2 × 100 mL). The solid was dried on the sinter under vacuum for 30 min, then transferred to a 250 mL round-bottomed flask and dried in vacuo to leave the solid precipitate. The solid was washed with toluene (2 × 100 mL) and dried on the sinter under vacuum for 30 min until a free-flowing powder was obtained. The crude product (typically 22–24 g) was transferred to a 500 mL round-bottomed flask containing a stirrer bar, followed by toluene (100 mL). A 10 mL Dean–Stark trap filled with toluene and a reflux condenser were fitted to the flask, and the suspension heated to reflux (180 °C DrySyn® temperature). Once at reflux, further 10 mL portions of toluene were added down the condenser, waiting ~5 min between portions, until a clear solution was obtained (~4–6 portions were typically needed). The reflux was maintained until no further H2O was observed condensing into the trap (typically 30 min were sufficient). The flask was removed from the heating block and allowed to cool slowly to r.t., over which time a precipitate formed. Once cooled, the flask was further cooled to ~10 °C in a NaCl/ice/water bath for 30 min. The solid precipitate was recovered by vacuum filtration on a 1.0 L sintered glass filter funnel (porosity 3), and washed with further portions of toluene (2 × 100 mL). The solid was dried on the sinter under vacuum for 30 min, then transferred to a 250 mL round-bottomed flask and dried in vacuo to constant weight. This gave title compound (R)-3 as fluffy white crystals; yield: 14.52 g (53.71 mmol, 77%); mp 163–164 °C (toluene). Analysis: calcd for C15H14N2OS: C, 66.64; H, 5.22; N, 10.36. Found: C, 66.49; H, 5.08; N, 10.27. Optical rotations were measured on an Electrothermal 9100 melting point apparatus. GC analyses were obtained on a Shimadzu GC-2025, with He as the carrier gas in split injection mode at constant linear velocity. GC analyses were performed on a Shimadzu GC-2025, with He as the carrier gas in split injection mode at constant linear velocity. All spectroscopic data were identical to those of (R)-3 using (S)-2-phenylglycinol ([S]-2; 10.00 g, 72.80 mmol) gave the title compound ([S]-3) as fluffy white crystals after a second recrystallisation; yield: 13.91 g (51.45 mmol, 74%); mp 158–159 °C (toluene); [α]20 D +101.7 (c 1.01, MeOH). GC analyses were obtained on a Shimadzu GC-2025, with He as the carrier gas in split injection mode at constant linear velocity. All spectroscopic data were identical to those of (R)-3. A 1.0 L round-bottomed flask containing a stirrer bar was charged with anhydrous CH2Cl2 (460 mL, 0.1 M) followed by (R)-3 ([21]D 12.42 g, 45.94 mmol, 1.0 equiv) with stirring and fitted with a suba seal and exit needle (19 gauge), Et3N (25.6 mL, 183.8 mmol, 4.0 equiv) was added via syringe and the suspension cooled in an ice/water bath. After 10 min, MsCl (4.62 mL, 59.72 mmol, 1.3 equiv) was added dropwise over ca. 5 min, during which time the suspension dissolved to give a pale yellow solution. The ice/water bath was removed and the reaction mixture stirred for 15 min. The reaction was checked by TLC ([CH2Cl2–Et2O 1:1; UV365/KMnO4; Rf (R)-3 = 0.28; 6 = 0.61; 7 = 0.81] and a further portion of MsCl (0.36 mL, 4.6 mmol, 0.1 equiv) added if (R)-3 remained (stir 15 min, re-check TLC, and repeat if necessary). It is crucial that all (R)-3 was consumed. Once complete consumption of
(R)-3 was observed, i-PrOH (9.0 mL) was added and the subea seal replaced by a reflux condenser. The reaction mixture was heated to reflux (50 °C DrySyn® temperature) overnight (ca. 18 h) and checked for completion by 1H NMR analysis. The reaction was quenched with aq 1 M NaOH (200 mL) and the biphasic mixture stirred vigorously for 30 min. The layers were separated and the aqueous layer extracted with CH2Cl2 (100 mL). The combined organics were washed with brine (100 mL), dried (MgSO4), filtered, and concentrated in vacuo. The crude residue was azeotroped with toluene (3 × 50 mL) to remove most of the residual Et3N. The residue was triturated with Et2O (200 mL with sonication and the liquid collected by decantation by a plug of cotton wool. This was repeated with two further portions of Et2O (100 mL) filtering the final portion through the cotton wool. The filter cake was rinsed with a further portion of Et2O (50 mL). The combined ethereal washings were extracted with aq 1 M HCl (5 × 50 mL) and the combined aqueous layers basified with aq 2 M NaOH (pH >14) at which point a cloudy white precipitate formed. The aqueous layer was extracted with Et2O (3 × 100 mL) and the combined organics washed with brine (100 mL). The organic layer was treated with activated charcoal (~5 g) and dried (MgSO4). The suspension was filtered through a pad of Celite, rinsing with further portions of Et2O (2 × 100 mL). The solvent was removed in vacuo to leave the crude product, which was purified by flash chromatography on silica gel (CH2Cl2) to leave (±)-BTM; (±)-4 as a colourless foam; yield: 661 mg (1.55 mmol, 84%); [α]20 +31.8 (c 1.0, CHCl3).

IR (film): 3028, 2936, 1636, 1580, 1454, 1352, 1244, 1173, 1157, 1034, 961 cm⁻¹.

1H NMR (500 MHz, CDCl3): δ = 2.97 (3 H, s, OSO2CH3), 3.67 (3 H, m, NSO2CH3), 4.33–4.52 (2 H, m, C(2)H) and 1H17F(4)H, 4.51 (5 H, dd, J = 8.4, 7.2 Hz, C(1)H17F(4)H, 7.12 (1 H, dd, J = 7.6, 1.1 Hz, ArH), 7.20–7.27 (2 H, m, Ar), 7.30–7.40 (3 H, m, and p-PhH), 7.43–7.46 (2 H, m, o-PhH), 7.98 (1 H, dd, J = 8.5, 0.7 Hz, ArH).

13C{1H} NMR (126 MHz, CDCl3): δ = 52.6, 75.5, 108.6, 121.6, 123.3, 126.6, 126.7, 127.5, 127.7, 128.9, 137.2, 143.1, 166.8.


IR (film): 3028, 2936, 1636, 1580, 1454, 1352, 1244, 1173, 1157, 1034, 961 cm⁻¹.

1H NMR (400 MHz, CDCl3): δ = 2.97 (3 H, s, OSO2CH3), 3.67 (3 H, m, NSO2CH3), 4.33–4.52 (2 H, m, C(2)H) and 1H17F(4)H, 4.51 (5 H, dd, J = 8.4, 7.2 Hz, C(1)H17F(4)H, 7.12 (1 H, dd, J = 7.6, 1.1 Hz, ArH), 7.20–7.27 (2 H, m, Ar), 7.30–7.40 (3 H, m, and p-PhH), 7.43–7.46 (2 H, m, o-PhH), 7.98 (1 H, dd, J = 8.5, 0.7 Hz, ArH).

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1H and 13C NMR data were consistent with literature values.2

(5)-(–)-2-Phenyl-2,3-dihydrobenzo[d]imidazo[2,1-b]thiazole [(5)-(–)-BTM; (5)-(–)-4] An identical procedure to that outlined for (R)-(–)-BTM 4 using (S)-3 (13.15 g, 48.64 mmol), MsCl (4.89 mL, 63.23 mmol), and Et3N (27.1 mL, 194.6 mmol) in anhydrous CH2Cl2 (490 mL) gave (S)-(–)-BTM (5)-(–)-4 in two crops as white crystals; yield: 8.96 g (35.51 mmol, 73%); mp 94–95 °C.

1H NMR spectrum was identical to that prepared by the optimised route.

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**Supporting Information**

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(17) From TCI-UK Ltd.: (R)-BTM P/N: B3296, £ 94.20/1 g; (S)-BTM P/N: B3549, £ 83.45/1 g (accessed 22/08/2014).

(18) Racemic 2-phenylglycinol is prohibitively expensive and was obtained by mixing equal amounts of the commercially available (R)- and (S)-2-phenylglycinols.

(19) See experimental for details.

(20) The site of N-mesylation was determined by $^1$H$^{15}$N HMBC spectroscopy, supported by DFT calculations for the $^{15}$N chemical shifts. See Supporting Information for details.

(21) Birman states that the addition of MeOH serves to quench additional MsCl, see ref. 1.

(22) MeOH was replaced with i-PrOH as it was considered that the by-product i-PrOMs from reaction with sulfene would be less reactive as an alkylating agent than MeOMs, thereby reducing polymeric impurities.


(24) GC analysis indicated >95% conversion of 2-chlorobenzothiazole against o-dichlorobenzene as the internal standard. Approximately 200 μL samples of the reaction mixture at t = 0 and 24 h were taken and diluted to ~1.5 mL in CH₂Cl₂ for analysis.