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## Enantioselective Total Synthesis of (—)-Pironetin: Iterative Aldol Reactions of Thiazolidinethiones

Michael T. Crimmins\* and Anne-Marie R. Dechert

Kenan and Venable Laboratories of Chemistry, University of North Carolina at Chapel Hill, Chapel Hill, North Carolina 27599

crimmins@email.unc.edu

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## ABSTRACT

The enantioselective total synthesis of pironetin has been achieved in 11 steps from known aldehyde 2. The synthesis relies on the formation of 5 out of 6 stereocenters through titanium mediated iterative aldol reactions. Key steps in this synthesis include an acetal aldol reaction to establish the stereochemistry at C8 and C9, an acetate aldol reaction, and "Evans" syn aldol reaction.

In 1994, the Kobayashi and Yoshida groups independently isolated the  $\alpha,\beta$ -unsaturated lactone (-)-pironetin **1** from the fermentation broth of *Streptomyces prunicolor* PA-48153 and *Streptomyces sp.* NK 10958.<sup>1</sup> (-)-Pironetin, originally identified as a novel plant growth inhibitor with immunosupressant activity, <sup>1b</sup> was later shown to possess antiproliferative activity against several tumor cell lines.<sup>2</sup> Further investigation established that (-)-pironetin has a unique mode of action compared to other tubulin binding agents, in that it covalently binds to the  $\alpha$  subunit of tubulin.<sup>3</sup>

(—)-Pironetin consists of an  $\alpha,\beta$ -unsaturated  $\delta$  lactone possessing a linear alkyl chain containing four contiguous stereogenic centers and a *trans*-olefin. To date, several other syntheses of (—)-pironetin and its derivatives have been reported.<sup>4</sup>

Most recently, Cossy reported an approach to (—)-pironetin utilizing enantioselecive crotylation and allylation methods, <sup>4m</sup> Enders and co-workers have disclosed a convergent total synthesis relying on RAMP/SAMP hydrazone methodology, <sup>4l</sup> and Nelson and co-workers have completed a total synthesis employing acyl halide-aldehyde cyclocondensations to construct the polypropionate units present in pironetin. <sup>4k</sup>

We have previously demonstrated the ability to execute iterative propionate aldol reactions for the synthesis of complex polypropionates in the context of the completion of a formal synthesis of deoxyerythronolide B.<sup>5</sup> A key goal in our designed approach to (—)-pironetin was to take

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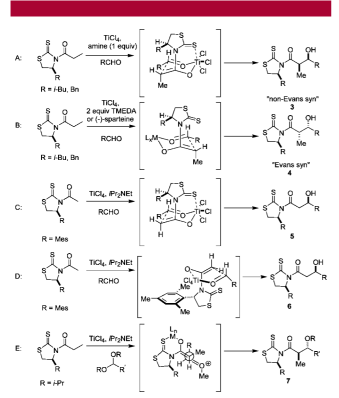
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advantage of an iterative aldol sequence that would exploit not only the syn aldol variant, but also other variations of the titanium tetrachloride mediated aldol additions of *N*-acylthiazolidinethiones to aldehydes and acetals. The demonstration that different types of polyketide frameworks found in natural products could be accessible through this technology was seen as an important goal of the synthesis.



**Figure 1.** Proposed transition states to access (A) "non-Evans" syn aldol adducts, (B) "Evans" syn aldol adducts, (C and D)  $\beta$ -hydroxy aldol adducts, and (E) anti  $\beta$ -alkoxy- $\alpha$ -methyl aldol adducts.

The use of chlorotitanium enolates of thiazolidinethione chiral auxiliaries allows access to either "Evans" syn and "non-Evans" syn aldol adducts in high diastereosectivity simply by changing the stoichiometry and nature of the amine base employed. This interesting tunability can be rationalized by a highly ordered transition state, in which the sulfur can either coordinate to the titanium metal center leading to "non-Evans" syn aldol adducts (Figure 1A), or in the case when a strongly coordinating amine ligand is used, inhibit the thiocarbonyl from chelating to the metal center resulting in "Evans syn" aldol adducts (Figure 1B).

Access to  $\beta$ -hydroxyl carbonyl subunits such as **5**, has proven more challenging with regard to obtaining high levels of diastereoselectivity due to the lack of substitution at the  $\alpha$  carbon of the enolate, however, we recently reported the use of the mesityl-substituted thiazoldinethione chiral auxiliary to access the  $\beta$ -hydroxyl carbonyl subunit in an efficient

and highly diastereoselective reaction. The high diastereoselectivity is thought to arise from a highly ordered chairlike transition state (Figure 1C), though a nonchelate boat-like transition state cannot be ruled out (Figure 1D).<sup>7</sup>

Direct access to the anti  $\beta$ -hydroxy aldol adducts is not currently possible using the chlorotitanium enolates. Several alternative approaches have been developed to access the β-hydroxy anti aldol adducts.<sup>8</sup> Evans and co-workers have developed an anti aldol reaction employing both oxazolidinones and thiazolidinethiones through the use of magnesium enolates. 8a,b Although aromatic aldehydes give highly diastereoselective aldol reactions in high yields using this method, aliphatic aldehydes give significantly lower conversion. A viable alternative to this method is the acetal aldol, developed by Urpi and co-workers, allowing access to anti  $\beta$ -alkoxyα-methyl aldol adducts.9 The high diasteroselectivity associated with the acetal aldol is thought to arise through an open transition state, in which the oxacarbenium ion is attacked from the less hindered face of the chelated Z enolate, in an antiperiplanar arrangement (Figure 1E).9

The Urpi anti acetal aldol seemed ideally suited to access the anti subunit at C8-9 of pironetin since a methoxy group, rather than a free hydroxyl group, resides at C9. Our strategy

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to access pironetin thus relies on the formation of 5 out of the 6 stereocenters through titanium tetrachloride mediated iterative aldol reactions<sup>5</sup> utilizing the thiazolidinethione chiral auxiliary to execute an anti propionate acetal aldol, an acetate aldol and a syn propionate aldol. Our retrosynthetic analysis is outlined in Scheme 1. We envisioned formation of the Z-enoate of **1** through a modified Horner-Emmons reaction <sup>10</sup> with aldehyde 8. Aldehyde 8 could be synthesized via an "Evans" syn aldol reaction with aldehyde 10 and thiazoldidinethione 9. Aldehyde 10 could be accessed via a highly diastereoselective acetate aldol reaction between mesitylsubstituted thiazolidinethione 11 and aldehyde 12. Aldehyde 12 would result from a highly diastereoselective acetal aldol reaction between a chiral dimethylacetal 14 and propionate 13. The stereocenter at C-10 would be set using an asymmetric alkylation, originally developed by Evans.<sup>11</sup>

The synthesis commenced with treatment of known aldehyde  $2^{12}$  with MeOH and p-TsOH to provide acetal 14. An acetal aldol reaction with the resultant dimethyl acetal 14 and propionate 13 according to the conditions described

by Urpi,  $^9$  provided the methylated aldol adduct **15** in a 64% yield with 98:2 dr. The reaction was optimized by employing an excess of the enolate (2 equiv), and using SnCl<sub>4</sub>, as opposed to BF<sub>3</sub>-OEt<sub>2</sub>, as the Lewis acid. The stereochemical outcome is further reinforced by the presence of the alpha stereocenter of the acetal resulting in Felkin control.  $^{96}$ 

Reductive cleavage<sup>12</sup> of the chiral auxiliary afforded aldehyde 12, which was subjected to an acetate aldol reaction<sup>7</sup> with thiazolidinethione **11**to afford alcohol **16** in 88% yield and 95:5 dr. An excess of the enolate (1.5 equiv) was necessary to achieve complete conversion of the aldehyde to the aldol adduct. Protection of alcohol 16 as its triethylsilyl ether delivered the thiazolidinethione 17. Reductive removal of the auxiliary with diisobutylaluminum hydride furnished aldehyde 10. Aldehyde 10 was then subjected to "Evans" syn aldol reaction conditions<sup>6</sup> employing an excess of the enolate of thione 9, affording aldol adduct 18 in >20:1 dr and a 65% yield. Silylation of the resulting alcohol provided triethylsilyl ether 19, which was then exposed to diisobutylaluminum hydride to effect reductive cleavage of the chiral auxiliary affording aldehyde 8. Thus, three iterative aldol reactions (8 steps overall) allowed the incorporation of 5 of the stereocenters of pironetin affording aldehyde 8.

Aldehyde **8** was treated with excess phosphonate **20**, to effect a modified Horner-Emmons reaction to access  $\alpha, \beta$ -

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unsaturated ester **21** as a 10:1 mixture of E/Z-isomers. Exposure of **21** to PPTS in 10:1  $CH_2Cl_2/MeOH$  provided only the unprotected diol, however, upon heating ester **21** with PPTS in 10:1 benzene/MeOH, both protecting groups were removed and lactonization was induced to furnish (-)-pironetin **1** in 63% yield (Scheme 2). Synthetic **1** was identical in all aspects to the natural product.

In summary, the enantioselective total synthesis of (—)-pironetin has been completed in 11 steps from previously prepared aldehyde **2** with an overall yield of 12.5%. In addition, the versatility of chlorotitanium mediated asymmetric aldol reactions was demonstrated through a sequence

of steps to rapidly construct pironetin in a highly stereoselective fashion.

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**Supporting Information Available:** Experimental procedures and copies of <sup>1</sup>H and <sup>13</sup>C NMR spectra. This material is available free of charge via the Internet at http://pubs.acs.org. OL9003228

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