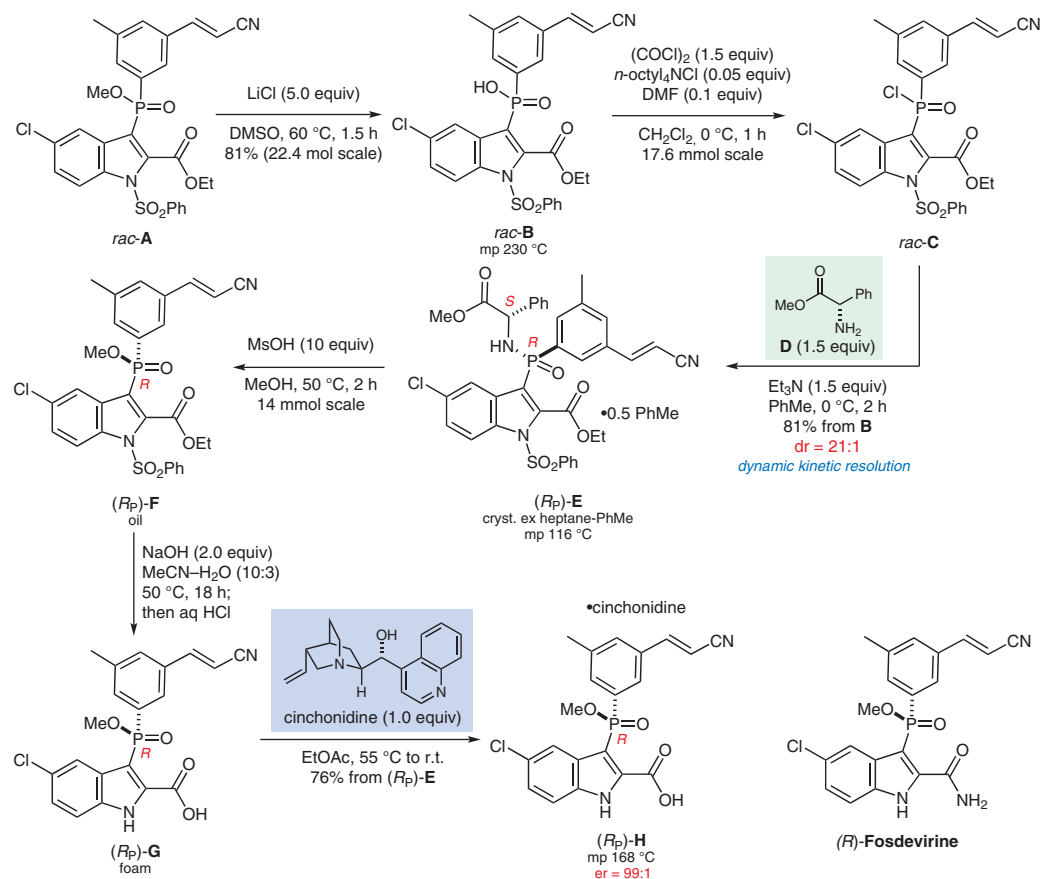


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A Highly Diastereoselective Chloride-Mediated Dynamic Kinetic Resolution at Phosphorus On-Route To A Key Intermediate In the Synthesis Of GSK2248761A

Tetrahedron Lett. **2018**, *59*, 2154–2156.

Synthesis of Fosdevirine



Significance: Fosdevirine (GSK2248761A) is a non-nucleoside reverse transcriptase inhibitor that was of interest for the treatment of HIV. A recent process-scale synthesis of the (*R*)-enantiomer was achieved by a late-stage classical resolution of racemic phosphinate **G** using cinchonidine as the resolving agent (*Org. Process Res. Dev.* **2018**, *22*, 200). A new route to key intermediate (*R_p*)-**G** in the synthesis of (*R*)-fosdevirine features a highly diastereoselective dynamic kinetic resolution in the reaction of the phosphinoyl chloride **rac-C** with methyl (*S*)-phenylglycinate (**D**).

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Comment: The key step in the dynamic kinetic resolution is the rapid racemization of phosphinoyl chloride **rac-C** by nucleophilic attack at the phosphorus atom by chloride ions followed by diastereoselective reaction of one of the phosphinoyl chloride enantiomers with the methyl (*S*)-phenylglycinate. Fifteen chiral amines were screened in the DKR reaction with best results (dr = 21:1) being obtained with methyl (*S*)-phenylglycinate. The overall yield of (*R_p*)-**H** from **A** was 50% (10 g scale).

Category

Synthesis of Natural Products and Potential Drugs

Key words

Fosdevirine

GSK2248761A

non-nucleoside reverse transcriptase inhibitor

dynamic kinetic resolution

phosphinate esters

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of the month

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