

Corrigendum

The authors of this Communication have recognized an error in Scheme 1. The correct Scheme 1 is shown here.

The text which refers to this Scheme (page 4597, left column) is also incorrect. It should read: "This observation suggests two alternative mechanisms for chain release coupled to isochromanone formation, to yield the putative intermediate deshydroxyajudazol B (3) (Scheme 1): TE-catalyzed attack of the C9 hydroxy group of the chain onto the acyl ester bond to give the free ten-membered ring lactone, followed by C2–C7 aldol addition and ring I aromatization; or b) aldol addition/aromatization to form ring I while the intermediate remains tethered to the TE, followed by TE-catalyzed lactonization and chain release to yield ring II."

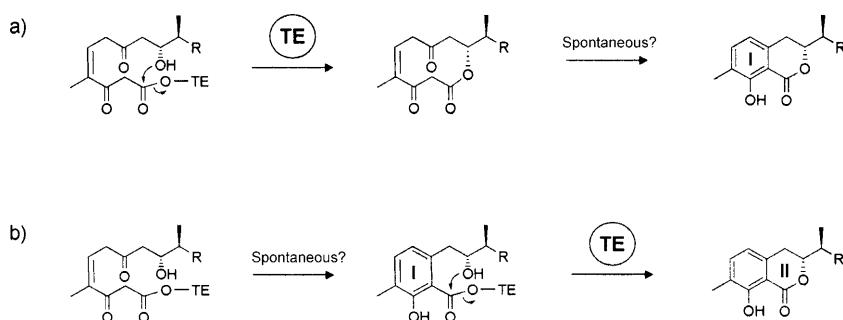
The authors would like to point out that this error does not affect the interpretation of data in the manuscript.

Production of the Antifungal
Isochromanone Ajudazols A and B in
Chondromyces crocatus Cm c5:
Biosynthetic Machinery and
Cytochrome P450 Modifications

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Scheme 1. Proposed mechanism for formation of the isochromanone ring system and chain release. a) The TE catalyzes lactone ring formation, which is followed by aldol addition and aromatization of ring I. b) Aldol addition and aromatization occur to generate ring I, followed by TE-catalyzed lactonization and chain release to afford ring II.