

Enantioselective Carbonyl Reverse Prenylation from the Alcohol or Aldehyde Oxidation Level Employing 1,1-Dimethylallene as the Prenyl Donor [J. Am. Chem. Soc. 2009, 131, 6916-6917]. Soo Bong Han, In Su Kim, Hoon Han, and Michael J. Krische*

In this report, the absolute stereochemical assignment of the reaction products was based upon correlation of HPLC data for adduct 4a to material prepared by Nakajima. However, upon use of this method to construct a known bryostatin substructure, 2a the opposite diastereoselectivity was observed. Additionally, in tert-prenylations of isatin, 2b the opposite enantioselectivity was observed. On the basis of these observations, as well as the comparison of adduct 4a to optical rotation data reported by Loh^{3a} and Denmark, ^{3b} we now report that our initial assignment of absolute stereochemistry was incorrect. That is, the (R)-enantiomers of the products are formed using the catalyst modified by (S)-SEGPHOS. This revision in absolute stereochemical assignment extends to recently reported tert-prenylations of furan methanols and furfurals. 2c Notably, the iridium-catalyzed *tert*-prenylations exhibit enantioselectivity opposite to that of corresponding allylations^{2d,e} and crotylations.^{2f}

Literature Cited

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- Han, S. B.; Krische, M. J. J. Am. Chem. Soc. 2009, 131, 2514.
 (3) (a) Loh, T.-P.; Zhou, J.-R.; Yin, Z. Org. Lett. 1999, 11, 1855. (b) Denmark, S. E.; Fu, J. J. Am. Chem. Soc. 2001, 123, 9488.

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10.1021/ja105736a Published on Web 08/17/2010 A Sulfido-Bridged Diiron(II) Compound and Its Reactions with Nitrogenase-Relevant Substrates [J. Am. Chem. Soc. 2004, 126, 4522-4523]. Javier Vela, Sebastian Stoian, Christine J. Flaschenriem, Eckard Münck,* and Patrick L. Holland*

Page 4522. The zero-field Mössbauer parameters determined for complex 1 were misreported. Therefore, in the second paragraph, fifth sentence, the statement " $\Delta E_{\rm O} = 0.58$ mm/s and isomer shift $\delta = 0.86$ mm/s" should read " $\Delta E_0 = 0.86$ mm/s and isomer shift $\delta = 0.58$ mm/s".

Supporting Information, page S-12. In the second sentence of the caption of Figure S5, " $\Delta E_{\rm O} = 0.58$ mm/s and $\delta = 0.86$ mm/s" should read " $\Delta E_{\rm O} = 0.86$ mm/s and $\delta = 0.58$ mm/s".

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Energetics of Phosphate Binding to Ammonium and Guanidinium Containing Metallo-Receptors in Water [J. Am. Chem. Soc. 2003, 125, 14807-14815]. Suzanne L. Tobey and Eric V. Anslyn*

Page 14812. In Table 3, first row, fourth column, the value for the enthalpy of binding 1 and HPO₄²⁻ should be ΔH° = -0.6 kcal/mol, not +0.6.

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