

# Additions and Corrections

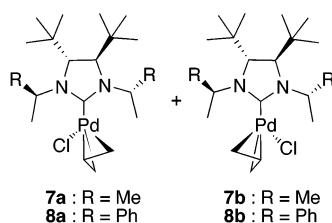
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**Sylvain Roland,\* Max Audouin and Pierre Mangeney:** Synthesis of Chiral (Diaminocarbene)Pd-(allyl)Cl Complexes

Page 3076 and table of contents. In Figure 3 and the structure in the table of contents entry, the absolute configuration of the carbons bearing the *tert*-butyl groups should be *R* for all the complexes. There is no inversion of configuration on the chiral backbone. Figure 3 should appear as it does below.

Page 3076. With an achiral ligand (allyl)Pd(L)X complexes are already chiral, even if all distances between the allyl carbons and the Pd are identical. Thus, with an enantiopure ligand two diastereomers are effectively formed.

Pages 3075 and 3076. Concerning ref 5, chiral (NHC)-Pd(allyl) complexes have also been used in asymmetric catalysis by M. Sigman (kinetic resolution of alcohols): Jensen, D. R.; Sigman, M. S. *Org. Lett.* **2003**, 5, 63. This paper should have been cited. We thank Matt Sigman and Klaus Pörschke for these corrections.



**Figure 3.**

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