

# Additions and Corrections

2001, Volume 20

**Adam H. Eisenberg, Felicia M. Dixon, Chad A. Mirkin,\* Charlotte L. Stern, Christopher D. Incarvito, and Arnold L. Rheingold:** Binuclear Palladium Macrocycles Synthesized via the Weak-Link Approach.

Pages 2052–2058. The reported distances and angles for structure **8** are incorrect. Due to a recently discovered error in the software, the bond distances and angles were generated from a previous refinement. One distance published in the article (correct distance of N2–C37 = 1.160(6) Å) and several of the distances and angles in the Supporting Information are significantly different. The updated data can be found in the Cambridge Structural Database with the CSD REFCODE of MIKNID.

OM020713L

10.1021/om020713l

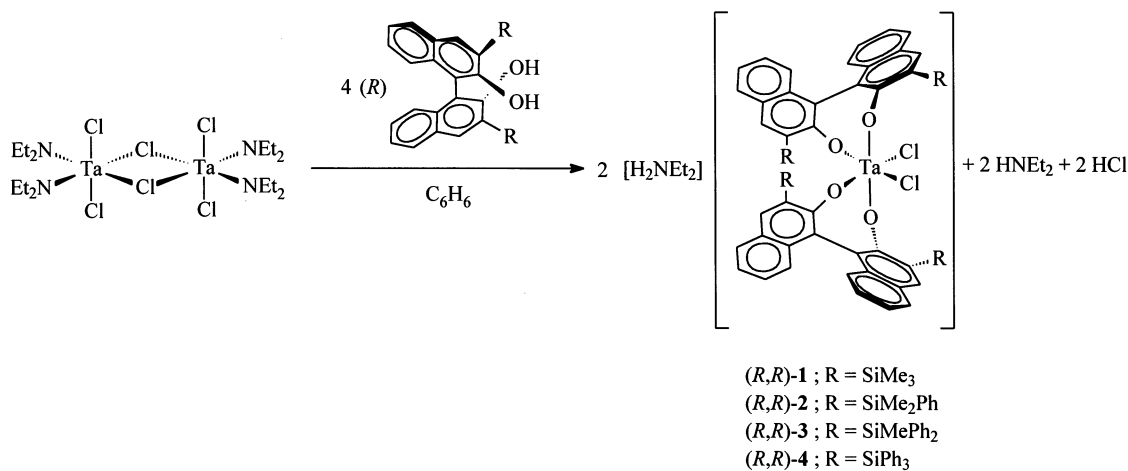
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2002, Volume 21

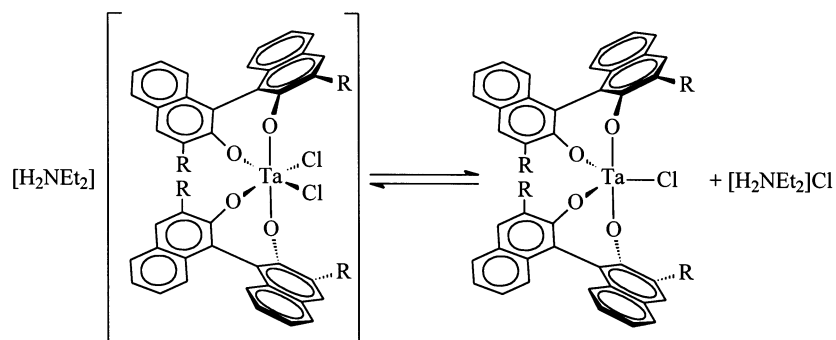
**Charles S. Weinert,\* Phillip E. Fanwick, and Ian P. Rothwell\*:** Isolation and Chemistry of Tantalum(V) Compounds Containing Two Resolved 3,3'-Disubstituted-1,1'-bi-2,2'-Naphthoxide Ligands

Pages 484–490. The structures of the 3,3'-dialkyl-1,1'-bi-2,2'-naphthoxido tantalum(V) complexes shown in Schemes 1–4 do not accurately represent the coordination environment about the metal centers. Both chelating ligands are in the  $\lambda$  configuration. Corrected schemes are shown below.

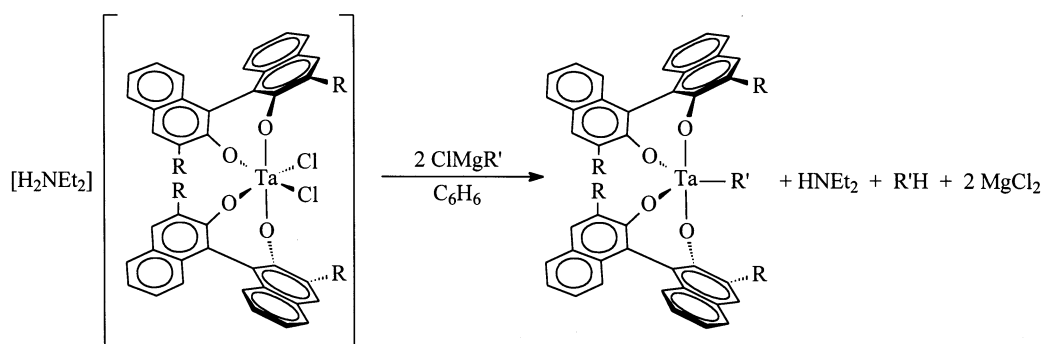
**Scheme 1**



Scheme 2



Scheme 3

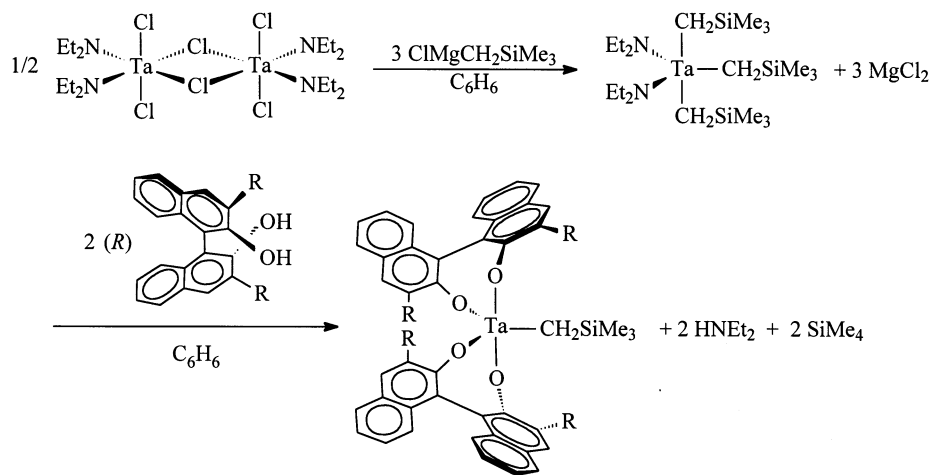


(*R, R*)-5 ; R = SiMe<sub>3</sub>; R' = CH<sub>2</sub>SiMe<sub>3</sub>

(*R, R*)-6 ; R = SiMe<sub>3</sub>; R' = CH<sub>2</sub>Ph

(*R, R*)-7 ; R = SiMe<sub>3</sub>; R' = *c*-C<sub>5</sub>H<sub>9</sub>

Scheme 4



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