

## Additions and Corrections

**Monomeric Cyclopentadienylnickel Methoxo and Amido Complexes: Synthesis, Characterization, Reactivity, and Use for Exploring the Relationship between M–X and H–X Bond Energies** [*J. Am. Chem. Soc.* **1997**, *119*, 12800–12814]. PATRICK L. HOLLAND, RICHARD A. ANDERSEN,\* ROBERT G. BERGMAN,\* JINKUN HUANG, AND STEVEN P. NOLAN

Page 12807: There is an important sign error in the first full paragraph in the second column. The first sentence of the paragraph should read: “The exchange between the amide ligand in **1a** and the cresolate ligand in **4** (eq 4) shows another equilibrium constant that is unexpectedly large...”. Later in the paragraph, in the sentence that begins, “An independent assessment...” the  $\Delta H(303\text{K})$  value should be changed to  $+9.4 \pm 0.2$  kcal/mol.

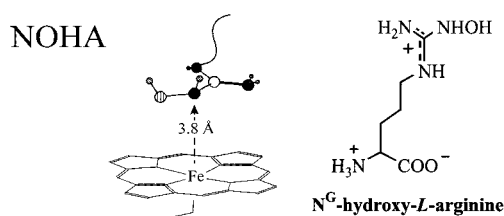
JA004657W

10.1021/ja004657w

Published on Web 09/06/2000

**ENDOR Studies of L-Arginine and N<sup>G</sup>-Hydroxy-L-Arginine Bound to All Three Holo-Nitric Oxide Synthase Isozymes** [*J. Am. Chem. Soc.* **2000**, *122*, 5405–5406]. DAVID L. TIERNEY, HUI HUANG, PAVEL MARTÁSEK, LINDA J. ROMAN, RICHARD B. SILVERMAN, BETTIE SUE SILER MASTERS, AND BRIAN M. HOFFMAN\*

Based on density functional theory calculations in Tantillo, D. J.; Fukuto, J. M.; Hoffman, B. M.; Silverman, R. B.; Houk, K. N. *J. Am. Chem. Soc.* **2000**, *122*, 536–537, the structure of N<sup>G</sup>-hydroxy-L-arginine in Scheme 1 should be drawn in the protonated form. Correspondingly, the structure of NOHA in Figure 1 should be as shown in the revised structure for NOHA below.



JA004658O

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