

ADDENDA AND ERRATA

Acta Cryst. (1994). C50, 470

Structures of 2,6-bis(aminomethyl)pyridine (bamp) complexes of Fe^{II}, Ni^{II}, Zn^{II}, Mn^{II}, Co^{III} and Cu^{II}. Erratum. By PIERRE BONHÔTE, MICHEL FERIGO, HELEN STOECKLI-EVANS and WERNER MARTY, *Institut de Chimie, Université de Neuchâtel, Avenue de Bellevaux 51, CH-2000 Neuchâtel, Switzerland*

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Abstract

An error in printing is corrected. In the paper by Bonhôte, Ferigo, Stoeckli-Evans & Marty [*Acta Cryst.* (1993). C49, 2102-2107] the *z* coordinate of the Cu atom of compound

(VI), chloro[2,6-bis(aminomethyl)pyridine]copper perchlorate, is given incorrectly as 1.00000. The correct value is 0.59397.

All relevant information is given in the *Abstract*.

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