

# Additions and Corrections

1968, Volume 7

**Stephen L. Lawton and Robert A. Jacobson:** The Crystal Structure of  $\alpha$ -Picolinium Nonabromoantimonate(V),  $(C_6H_5NH)_2Sb^VBr_9$ .

Page 2127. In Table II, atom Br<sub>2</sub>,  $B_{23}$  should be -1 (2).

1971, Volume 10

**Stephen L. Lawton, Robert A. Jacobson, and Robert S. Frye:** The Crystal Structure of Pyridinium Tetracosabromoantimon(III)triammon(V)ate,  $(C_6H_5NH)_6Sb^{III}Sb^V_3Br_{24}$ .

Page 701. In the Introduction, column 1, line 15, the temperature range should read 116-138°.

Page 704. In Table I, the value of the  $x$  coordinate for Sb(1) should be 0.22101, the standard deviation for the  $y$  coordinate of Sb(3) should be (19), and the footnote reference for the heading Group B should be f.

Page 706. In Table V three of the Sb-Br bond lengths *corrected for libration* are listed incorrectly and should read as follows: Sb(1)-Br(1), 2.545 (10), not 2.555 (10); Sb(3)-Br(10), 2.809 (10), not 2.801 (10); Sb(3)-Br(11), 2.801 (10), not 2.809 (10). The entries in Table VI for  $(C_6H_5NH)_6Sb_4Br_{24}$  are correct.

**Stephen L. Lawton and Robert A. Jacobson:** The Crystal

Structure of Quinuclidinium Dodecabromoantimon(III)antimon(V)ate-2-Dibromine,  $(C_7H_{15}NH)_4Sb^{III}Sb^VBr_{12} \cdot 2Br_2$ .

Page 711. In the heading of Table IV, the scaling of the observed and calculated structure factors should read: (in electrons  $\times 10$ ).

**James N. Francis and M. Frederick Hawthorne:** Synthesis and Properties of Cobalt Complexes Containing the Bidentate  $\pi$ -Bonding  $B_8C_2H_{10}^{4-}$  Ligand.

Page 863. The caption to Figure 1 should read: Schematic drawing of the (3,6)-dicarbacanastide ion with H atoms omitted.

**Melvin R. Churchill, Joan Cooke, James P. Fennessey, and John Wormald:** The Crystal Structure of Bis(imidotetramethyldithiophosphino-S,S)nickel(II), a Tetrahedral Complex with an NiS<sub>4</sub> core.

Page 1032. In Table I, the  $x$  coordinate of the nickel atom is given as 0.73097 (10); it should read 0.73907 (10). The  $y$  coordinate of S(3) is given as 0.48892 (37) and should read 0.48829 (37). All geometric information is based upon the correct atomic positions.

**Marvin L. Hackert, Robert A. Jacobson, and Timothy A. Keiderling:** The Crystal Structure of Tetraethylammonium Hexabromoantimonate(V),  $(C_2H_5)_4NSbBr_6$ .

Page 1075. In ref 9 the year should be 1971.