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Compounds with Transition Metal–Carbon π -Bonds
and Compounds of Groups 10 – 8 (Ni, Pd, Pt, Co, Rh, Ir,
Fe, Ru, Os)

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Volume 3:

Compounds of Groups 12 and 11 (Zn, Cd, Hg, Cu, Ag, Au)

3.1 Product Class 1: Organometallic Complexes of Zinc

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Volume 4: Compounds of Group 15 (As, Sb, Bi) and Silicon Com- pounds

4.4 Product Class 4: Silicon Compounds

4.4.46 Product Subclass 46: Siloles

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Volume 20:

Three Carbon—Heteroatom Bonds: Acid Halides; Carboxylic Acids and Acid Salts; Esters, and Lactones; Peroxy Acids and $R(CO)OX$ Compounds; $R(CO)X$, $X = S, Se, Te$

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