

Science of Synthesis

Knowledge Updates 2014/4

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Volume 6: Boron Compounds

6.1 Product Class 1: Boron Compounds

6.1.42 Product Subclass 42: N-Heterocyclic Carbene Borane Complexes

New

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 $X_2C=X$, CX_4**

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27.1.7 Sulfur Ylides

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