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Volume 24: Three Carbon—Heteroatom Bonds: Ketene Acetals and Yne—X Compounds

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Volume 31:

Arene—X (X = Hal, O, S, Se, Te, N, P)

31.5 Product Class 5: Phenols and Phenolates

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X—Ene—X (X = F, Cl, Br, I, O, S, Se, Te, N, P), Ene—Hal,
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