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Titolo	Metal-carbon bonds in enzymes and cofactors / / edited by Astrid Sigel, Helmut Sigel, and Roland K.O. Sigel
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	Coenzymes
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Nota di bibliografia	Includes bibliographical references and indexes.
Nota di contenuto	Organometallic chemistry of B12 coenzymes Cobalamin- and corrinoid-dependent enzymes Nickel-alkyl bond formation in the active site of methyl-coenzyme M reductase Nickel-carbon bonds in acetyl-coenzyme a synthases/carbon monoxide dehydrogenases Structure and function of [NiFe]-hydro-genases Carbon monoxide and cyanide ligands in the active site of [FeFe]-hydrogenases Carbon monoxide as intrinsic ligand to iron in the active site of [Fe]- hydrogenase Dual role of heme as cofactor and substrate in the biosynthesis of carbon monoxide Copper-carbon bonds in mechanistic and structural probing of proteins as well as in situations where copper is a catalytic or receptor site Interaction of cyanide with enzymes containing vanadium, manganese, non-heme iron, and zinc Reaction mechanism of the molybdenum hydroxylase xanthine oxidoreductase: evidence against the formation of intermediates having metal-carbon bonds.
Sommario/riassunto	The occurrence of a wide variety of metal-carbon bonds in living organisms, ranging from bacteria to humans, is only recently recognized. Of course, the historical examples are the B12 coenzymes containing cobalt-carbon bonds, but now such bonds are also known for nickel, iron, copper, and other transition metal ions. There is no

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