

Calculated tautomeric equilibria and X-ray structures of 2-substituted N-methoxy-9-methyl-9H-purin-6-amines

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Controlling chirogenic effects in porphyrin based supramolecular systems: theoretical analysis versus experimental observations

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Efficient defect-driven cation exchange beyond the nanoscale semiconductors toward antibacterial functionalization

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Solution-mediated inversion of SnSe to Sb2Se3 thin-films

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Supramolecular chirogenesis in a sterically hindered Porphyrin : a critical theoretical analysis

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