

Contents

II/20 Molecular Constants

Subvolume C: Non-linear Triatomic Molecules

Part 1: H₂O (HOH)

Part δ: D₂¹⁶O (D¹⁶OD), D₂¹⁷O (D¹⁷OD), D₂¹⁸O (D¹⁸OD), T₂¹⁶O (T¹⁶OT), T₂¹⁸O (T¹⁸OT)

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