

ELECTRONIC STATE OF CoH: A PROBLEM RELATED TO THE SIZE-INCONSISTENCY OF THE MR-SDCI METHOD AND THE MIXING OF $^3\Phi$ AND $^3\Pi$

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The ground state of CoH has been determined experimentally as the $^3\Phi$ by the far-infrared magnetic resonance study,^a and theoretically by *ab initio* methods.^b However, MR-SDCI and MR-SDCI + Q *ab initio* molecular orbital calculations predicted that the lowest electronic state is $^5\Phi$. The size-consistent MR-ACPF method correctly predicted that $^3\Phi$ should be the ground state. As is previously reported by Tanaka *et al.*^c on FeH, the controversial results in MR-SDCI may arise through truncation error inherent in the size-inconsistent MR-SDCI method.

Another difficulty is that we cannot clearly separate the Φ and Π states when CoH is treated under C_{2v} symmetry, not under its real symmetry of $C_{\infty v}$. The best results we obtained at the moment are: $r_e = 1.528 \text{ \AA}$, $B_e = 7.291 \text{ cm}^{-1}$, and $\omega_e = 2005 \text{ cm}^{-1}$ at the level of the MR-ACPF + relativistic correction using the Roos ANO (Co) and Dunning aug-cc-pVQZ (H) basis sets.

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