

THEORETICAL INVESTIGATION OF THE M^+-RG_2 (M = ALKALINE EARTH METAL; RG = RARE GAS) COMPLEXES

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Metal cation rare gas complexes provide an expectedly simple system with which to investigate intermolecular interactions. Despite this, we have previously found the M^+-RG (M = alkaline earth metal) complexes to very complicated systems, with the complexes of the heavier rare gases displaying surprisingly large degrees of chemical character.^{a,b,c} Here we extend these studies by examining the nature of these interactions with increasing degrees of solvation through investigating the M^+-RG_2 complexes using high level *ab initio* techniques. Intriguing trends in the geometries and dissociation energies of these complexes have been observed and are rationalized.

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