Chemical Bonding of Transition-metal Co13 Clusters with Graphene

T. Alonso-Lanza¹, A. Ayuela¹ and F. Aguilera-Granja²
¹Centro de Fisica de Materiales CFM-MPC CSIC-UPV/EHU, Donostia International Physics Center (DIPC), Departamento de Fisica de Materiales, Fac. de Quimicas, UPV-EHU, 20018 San Sebastian, Spain
²Instituto de Fisica, Universidad Autonoma de San Luis de Potosi, 78000 San Luis Potosi S.L.P., Mexico

tomas_alonso001@ehu.eus

Abstract
We carried out density functional calculation to study Co13 clusters on graphene [1]. We deposit several free isomers in different disposition respect to hexagonal lattice nodes, studying even the hcp 2d isomer recently obtained as the most stable one [2]. Surprisingly, Co13 clusters bonded to graphene prefer icosahedron-like structures where the low lying isomer is much distorted, because it is linked with more bonds than in previous works [3].

For any isomer the most stable position binds to graphene by the Co atoms that can lose electrons. We find that the charge transfers between graphene and clusters are small enough to conclude that the Co-graphene binding is not ionic-like but chemical. Besides, the same order of stability among the different isomers on doped graphene is well kept. These findings could also be of interest for magnetic clusters on graphenic nanostructures such as ribbons and nanotubes.

References

Figures