Search for Advanced Functional Materials concern with Eco-products

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Abstract
Graphenes have mono atomic-layer and honeycomb lattice of carbon materials in which it looks like a structure of physical peeling from graphite. It is a candidate for next generations of electronic devices, such as sensors, conductive films, catalysts and so on. For applications of these devices, we have interests in the feature of electronic structures of the graphenes. In this particular project, we focus upon the disability of Hydrogen evolution reactions (HER) under the conditions of the Oxygen reduction reaction (ORR) on the graphene materials. Additionally, we explained its unique affections for Nitrogen substitutions.

In this work using the Earth Simulator, we obtained the following results of the disability of HER on edged-graphene with a nitrogen-substitution:

1. For the comparisons of electron affinity between pristine and nitrogen substituted graphene, the armchair edged graphene shows decreasing affinity when nitrogen substitute in the edge - region, while zigzag-edged graphene shows increasing affinity when nitrogen substitute in edge-region.

2. For armchair-edged graphene ribbon, the adsorption energy of a proton on Nitrogen substituted graphene is much stronger than the adsorption energy of a proton on the pristine graphene. While for zigzag-edged graphene ribbon, their energy is weaker than the pristine graphene case.

Computational Details
In this work, we applied density functional theory (DFT) calculations with projector-augmented plane wave basis functions to edged-graphene. Our DFT studies used the Perdew-Burke-Ernzerhof (PBE) functional and exchange-correlation interactions under the generalized gradient approximations (GGA) frameworks. The wave-functions cutoff energy and the charge density cutoff energy were 30 and 240 Ry, respectively. The k-points mesh generation scheme used Monkhorst-Pack special point. All calculations of this study were carried out using the DFT program package of PHASE. The actual performance of the present version of PHASE showed 0.770 TFLOPS using 4 nodes of the Earth Simulator.
Keywords: Density functional theory, first-principles calculations, graphene, Oxygen reduction reaction, Hydrogen evolution reaction