

Electronic Supplementary Information for:

Metal-free Catalytic Reduction of 4-Nitrophenol to 4-Aminophenol by N-Doped Graphene**

Xiang-kai Kong, Zhi-yuan Sun, Min Chen, Chang-le Chen,* and Qian-wang Chen*

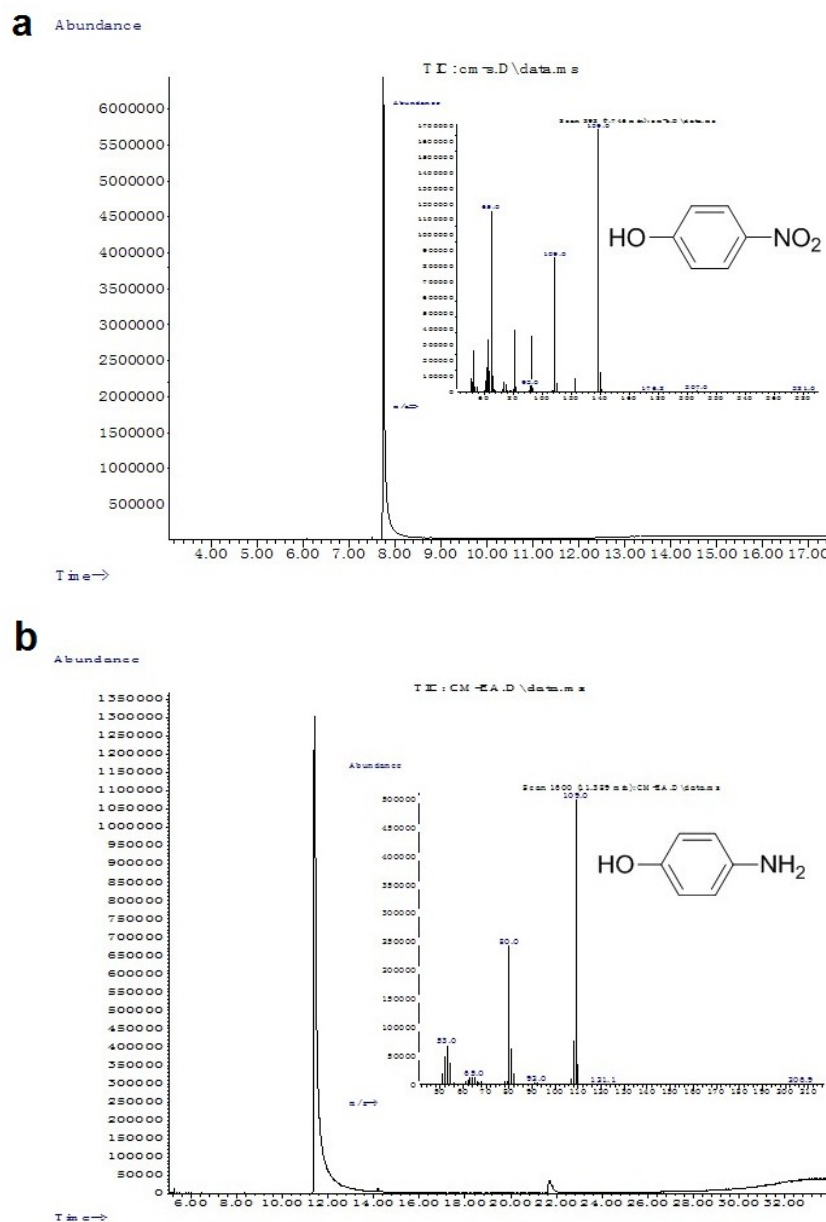


Figure S1. The GC-MS results of (a) the reactant Nip and (b) the Amp product after extracting the reaction mixture with ethyl acetate. (The small peak at retention time ca. 22min is unknown impurity, from the solvent or the small pieces of graphene. However, it is not Nip based on the retention time and the mass spectrum)

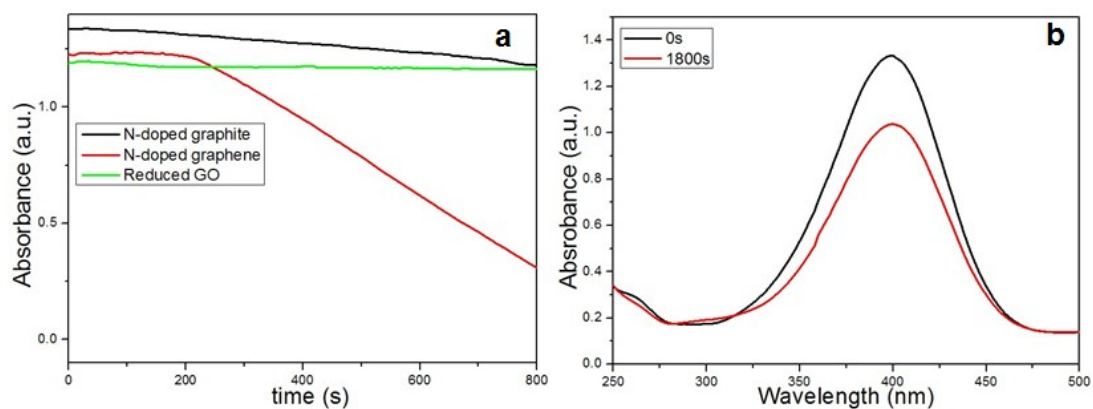


Figure S2. (a) The measured curves of absorbance at 400 nm vs. time for N-doped graphene, N-doped graphite and reduced GO; (b) The UV/Vis spectra of reduction Nip for N-doped graphite.

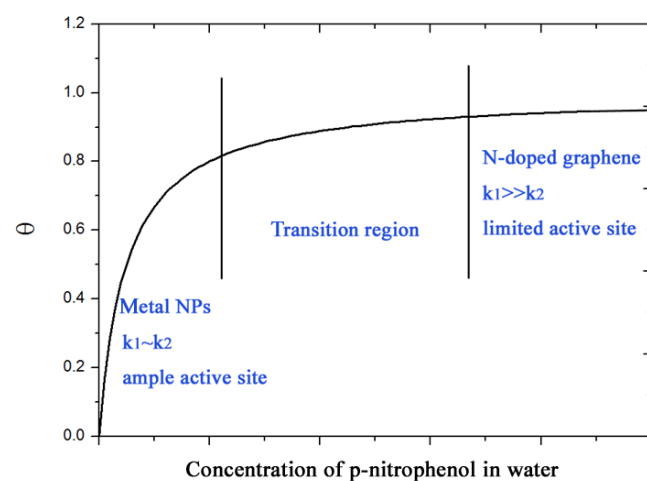


Figure S3. Diagram of Langmuir adsorption isotherm for metallic nanoparticles and N-doped graphene.

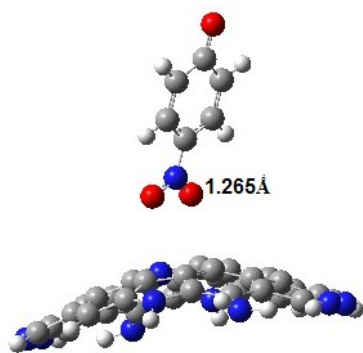


Figure S4. The optimized structure of Nip ion absorbed on the N-doped graphene sheet via its nitro group, and the large separated distance indicates a weak interaction at the interface. The N-O bond length has been marked directly.

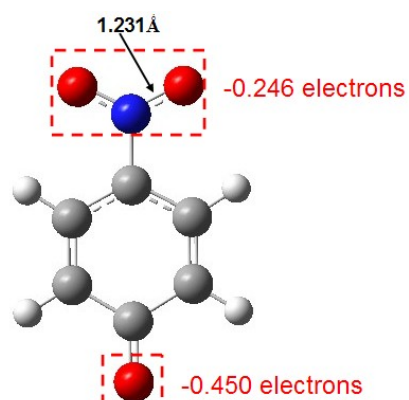


Figure S5. The calculated charge distributions for the nitro group and the O atom (adsorption site) of one free Nip ion, with the $\text{N}-\text{O}$ bond marked directly.

| Models | E _{ads} (Hartree) |
|---|----------------------------|
| N-doped graphene | -0.0665 |
| N-doped graphene via nitro group adsorption | -0.0282 |
| pristine graphene with center adsorption | 0.0006 |
| pristine graphene with edge adsorption | -0.0009 |
| graphene doped with pyridinic N atoms | -0.0082 |
| graphene doped with pyrrolic N atoms | -0.0173 |
| graphene doped with amine like N atoms | -0.0752 |
| graphene doped with graphitic N atoms | -0.0549 |

Table S1. The calculated Nip adsorption energies for these discussed graphene models.