



Cite this: *Phys. Chem. Chem. Phys.*,
2026, **28**, 980

Correction: Theoretical study of large-scale graphene on the Cu(111) surface using machine learning potential

Jingli Han, ^a Rubén Cabello, ^c Jordi Bonet Ruiz, ^c
 Alexandra Elena Plesu Popescu, ^c Sergi Dosta Parras, ^{*d}
 Camila Barreneche ^d and Yongpeng Yang ^{*b}

DOI: 10.1039/d5cp90209f

rsc.li/pccp

Correction for 'Theoretical study of large-scale graphene on the Cu(111) surface using machine learning potential' by Jingli Han *et al.*, *Phys. Chem. Chem. Phys.*, 2025, **27**, 17717–17729, <https://doi.org/10.1039/D5CP02042E>.

The authors regret that in the originating article, the affiliations of Yongpeng Yang and Jingli Han were incorrect. Their correct affiliation is Henan Institute of Advanced Technology, Zhengzhou University, Zhengzhou 450052, PR China, and School of Material and Chemical Engineering, Zhengzhou University of Light Industry, Zhengzhou, 450001, China, respectively.

The Royal Society of Chemistry apologises for these errors and any consequent inconvenience to authors and readers.

^a School of Material and Chemical Engineering, Zhengzhou University of Light Industry, Zhengzhou, 450001, China

^b Henan Institute of Advanced Technology, Zhengzhou University, Zhengzhou 450052, PR China. E-mail: ypyang2017@zzu.edu.cn

^c Department of Chemical Engineering and Analytical Chemistry, Faculty of Chemistry, University of Barcelona, Barcelona, 08028, Spain

^d Department of Materials Science and Physical Chemistry, Faculty of Chemistry, University of Barcelona, Barcelona, 08028, Spain. E-mail: sdosta@ub.edu

