Atomic and electronic structure of Co induced identical clusters grown on Si(111)-(7×7) template

M. A. K. Zilani, Y. Y. Sun, H. Xu, T. Liu, Liu Lei, Y. P. Feng, X.-S. Wang and A. T. S. Wee*

Department of Physics, National University of Singapore, Kent Ridge, Singapore 119260, Republic of Singapore

ABSTRACT

Cobalt-induced identical clusters have been grown on Si(111)-(7×7). In situ scanning tunneling microscopy (STM) studies show that the clusters are accompanied by voids created by two missing adatoms. We explain the atomic structure of these clusters by first-principles total energy calculations. The electronic structure of the clusters are investigated by bias dependent STM, scanning tunneling spectroscopy and real-space multiple-scattering calculations. Spectroscopic studies on the clusters indicate a band gap of ~0.8 eV, suggesting localized nonmetallic behavior.

INTRODUCTION

The template nature of Si(111)-(7×7) has been frequently used to grow nano-dots of different elements with identical size and shape comprising fixed number of atoms i.e. identical clusters and, attracted much interest in recent days [1-2]. Such low dimensional structures with interesting quantum properties are important not only for fundamental scientific interest but also for wide variety of potential nanotechnology applications. We have studied the early stage interaction of reactive metal Co with Si(111)-(7×7) [3,4]. At lower coverages Co-induced identical clusters are formed on Si(111)-(7×7) at specific adsorption sites in the faulted half of the unit cell [3]. In situ STM studies identify the interaction sites of the clusters. Each cluster is closer to one side of the FHUC, breaking the local three-fold symmetry, in stark contrast to all previous group-III cluster systems where the symmetry is preserved on Si(111)-(7×7) [2]. Based on first-principles total energy calculations and experimental observations, we propose a model to explain the atomic structure and formation mechanism of these clusters. The electronic structure of these clusters has been unveiled by a combined study of bias dependent STM, scanning tunneling spectroscopy and real space multiple scattering calculation [5].

RESULTS AND DISCUSSION

A. Co induced identical cluster formation on Si(111)-(7×7)

- STMS: ~0.06 ML Co at temperature 230±20 °C. F and U for faulted and unfaulited triangular half.
- STM images
- (a) Top and side view: 3 Si atoms at topmost layer + 6 Co atoms at next layer
- (b) 3 liberated Si center adatoms

B. Two magic clusters at both halves of a single unit cell

- Magic cluster: Localized nonmetallic with a band gap ~0.8 eV
- Flat region around Fermi level for Si
- Two surface states at -1.7 V and 1.2 V,
- Dominant peak Co 3p
- For clean Si surface area:
  - -0.35 V: Completely filled rest atoms
  - -0.7 V: Completely filled rest atoms
  - -1.4 V: Corner hole and Si-Si backbonds around the adatoms
- A band gap of 0.8 eV opens near the Fermi energy. This is important for understanding their electrical, optical and magnetic properties. The STMS results suggest that the identical clusters are responsible for a local removal of the metallicity of Si(111)-(7×7) and a band gap of 0.8 eV opens near the Fermi energy. This bandgap opening suggests the gain of electronic energy as a driving force for the formation of the clusters.

C. Formation mechanism and atomic Structure

- Method:
  - Calculations are based on DFT and ultrasoft pseudopotential plane-wave method.
  - Vacuum layer ~30 Å, atomic slabs with 8 layers Si.
  - Total 4 Cl. 396 Si and 49 H atoms considered per unit cell.
  - Cutoff energy of 17.5 Ry
- Results:
  - Site X filled by a Co
  - Permanent vacancy left at Site X
- More evidence of cluster interaction site on 7×7 template
  - (a) Two oppositely-directed clusters are connected back to back
  - (b) Voids of the two oppositely directed clusters share a common axis

D. Calculation of STM images

- Empty
- Filled
- Comparison of calculated and experimental STM images

References:

E. Electronic structure

- (i) Bias Dependent STM
- (ii) STS Measurements on a magic cluster
- (iii) RSM5 calculations of element and momentum projected DOS

Conclusion

The atomic and electronic structure study of low-dimensional systems such as 3d ferromagnetic metal induced clusters is important for understanding their electrical, optical and magnetic properties. The STS results suggest that the identical clusters are responsible for a local removal of the metallicity of Si(111)-(7×7) and a band gap of 0.8 eV opens near the Fermi energy. This bandgap opening suggests the gain of electronic energy as a driving force for the formation of the clusters.

*Corresponding author: FAX 65 67761426
Email address: phyweets@nus.edu.sg