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Structure of germanene/Al(111): a two-layers surface alloy

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Supporting Information

1. Details of the minimization procedure used for exploring all configurations.

In order to determine the structure of the surface, we have tested all possible configurations involving as well Al and Ge atoms in a bilayer configuration on top of a Al(111) substrate. For the first layer (surface plane), we have explored any position of the atoms that respects the space group symmetries, assuming at least 3 Ge atoms and at most 9 atoms in the surface layer. We have assumed that the space group was the same as the one of the substrate, i.e. p3m1. Configurations are defined by the occupancy of 1a, 1b, 1c, 3d and 6e Wyckoff positions.¹ Within these assumptions, the number of different configurations for the surface

plane is 118. For the second layer (interfacial plane), we have started from the atomic positions of an Al(111) plane where at most three Al atoms are replaced with Ge atoms, also respecting the p3m1 symmetries. For an Al plane, there are three possible atomic configurations for the 9 atoms inside a (3×3) cell. They are drawn in Fig. S1a-c. The first one corresponds to atoms belonging to the Wyckoff positions 1*a*, 1*b*, 1*c*, and 6*e*. The two other configurations correspond to three groups of atoms in 3*d* positions.



Figure S1: (a-c) atomic configurations for Al(111) planes respecting the p3m1 symmetries. (d) Ge₄Al₄ surface plane. (e) Ge₈ surface plane. (f) Ge₂Al₇ interfacial plane. The (3×3) unit cell is drawn in red, Al atoms are drawn in brown, Ge atoms in green. Wyckoff positions are indicated by *a*, *b*, *c*, *d*, *e* letters. Mirror and glide planes are drawn in black continuous and dotted lines respectively. Triangles indicate 3-fold rotation axes.

The maximum relaxation allowed for atoms of the interface plane was $\pm 10\%$ of the unit cell size. The third layer was considered as a pure Al plane, where atoms could relax with a maximum amplitude of 5% of the unit cell size. Debye Waller factors were used for the description of atomic vibrations in directions parallel or perpendicular to the surface. Debye-Waller factors were set independent for each group of symmetrical atoms within the first two planes, and equal for atoms of the third plane.

For each configuration tested, the best fit of the structure factors has been obtained by exploring the space of free parameters (a scale factor, the atomic positions and Debye-Waller factors) using the genetic algorithm implemented in SciPy. The agreement was obtained by minimizing $\sum \left(\frac{F_{\text{th}}-F_{\text{exp}}}{\sigma_{\text{exp}}}\right)^2 + E$, where F_{exp} and F_{th} are the experimental and simulated structure factors, σ_{exp} the experimental uncertainties and where *E* is a dimensionless Lennard-Jones interaction energy between nearest-neighbors atoms. *E* is given by:

$$E = \sum_{\substack{i,j \\ d_{ij} < 1.1d_0}} A\left[\left(\frac{d_0}{d_{ij}} \right)^{12} - 2\left(\frac{d_0}{d_{ij}} \right)^6 + C \right]$$

where d_{ij} is the distance between atoms *i* and *j*. *A* and d_0 depends on the chemical nature of atoms *i* and *j* and are given in table S1. *C* is a constant term equal to $2 \times 1.1^{-6} - 1.1^{-12}$ used to avoid any discontinuity of the interaction at the cutoff value.

(<i>i</i> , <i>j</i>)	Α	d_0 (nm)
Al-Al	5	0.286
Ge-Ge	1	0.245
Al-Ge	2	0.266

Table S1: parameters for the Lennard-Jones interaction



Figure S2: Models of Ge layers on Al(111) relaxed by DFT. (a) pure Al, (b) Ge1Al8/Al9, (c) Ge3Al6/Al9, (d) Ge3Al6/Ge1Al8, (e) Ge4Al4/Al9, (f) Ge4Al4/Ge1Al8, (g) Ge4Al4/Ge2Al7, (h) Ge4Al4/Ge3Al6, (i) Ge5Al3/Al9, (j) Ge5Al3/Ge1Al8, (k) Ge5Al3/Ge2Al7, (l) Ge5Al4/Ge1Al8, (m) Ge8/Al9 (1H), (n) Ge8/Ge1Al8, (o) Ge8/Ge2Al7, (p) Ge8/Ge3Al6. The side views correspond to a projection along a plane 7° off from the $(11\overline{2})$ plane.