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Iuliacumite: a novel two-dimensional chemical short range order in a wurtzite single monolayer InAs$_{1-x}$Sb$_x$ shell on InAs nanowires

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A chemical short range order is found in single monolayer InAs$_{1-x}$Sb$_x$ shells, which inherit a wurtzite structure from the underlying InAs nanowire, instead of crystallizing in the energetically preferred zincblende structure. The chemical order is characterized by an ordering vector in $[0001]$ and an anti-ordering vector in $(\bar{1}1\bar{2}0)$ direction and arises from strong Sb-Sb repulsive interactions along the atomic chains in $(\bar{1}1\bar{2}0)$ direction.

The reduced dimensionality of semiconductor nanowires (NWs) offers the unique opportunity to grow materials with crystal structures, which are otherwise unstable. One of the most prominent examples is the growth of wurtzite (WZ) structure InAs NWs,$^4$ although stable InAs bulk material prefers the zincblende (ZB) structure. In contrast, other III-V materials, notably InAs$_{1-x}$Sb$_x$, prefer to keep their ZB bulk structure even in NWs or nanostructures with reduced dimensionality.$^{[2,3]}$ However, lateral overgrowth of these ternary III-V semiconductor alloys on sidewall facets of WZ structure III-V nanowires offers the prospect to nevertheless obtain reliably a WZ structure shell, despite being unstable. Thereby new polytype structures of ternary III-V semiconductor alloys can be achieved, offering additional degrees of freedom for adjusting the band structure in, e.g., core-shell nanowires.

In addition of controlling the polytype structure, controlling ordering of the different chemical species in ternary III-V semiconductor alloys offers further means to tune the electrical properties. This has been amply demonstrated for ZB structure III-V semiconductors, where a variety of different chemical ordering structures were identified (i.e. chalcopyrite, CuPt, CuAu, famatinite, triple period, and lazarevicite).$^{[4–9]}$ In contrast, rather little is known about chemical ordering in WZ structure III-V semiconductors. Thus far most reports on chemical ordering refer to $c$ axis ordering and superlattices in WZ structure group III-nitride semiconductors.$^{[10–17]}$ Only during growth on pyramidal pits ordering along the pit’s semi-polar growth facets was observed.$^{[18]}$ In contrast to the limited experimentally observed ordering structures in WZ materials, theory predicts a large variety of possible ordered WZ structures, with ordering vectors not only in $c$ direction$^{[19]}$ $^{[20]},$ but without experimental verification yet.

Furthermore, thus far chemical ordering in III-V semiconductor alloys was identified in (quasi) three-dimensional materials, i.e. bulk or extended nanowire bulk-like sections. However, lateral overgrowth on NW sidewall facets offers the possibility to obtain monolayer thin two-dimensional ternary III-V semiconductor sheets. At present it is fully unclear if any chemical ordering is forming in such two-dimensional WZ-structure III-V semiconductors.

Therefore, we designed a two-dimensional single monolayer InAs$_{1-x}$Sb$_x$ WZ structure shell on sidewall facets of InAs nanowires and investigated the chemical ordering using atomically resolved scanning tunneling microscopy (STM). We identify the existence of a short-range chemical ordering in this WZ structure single monolayer shell. The new type of two-dimensional ordering, called Iuliacumite, is characterized by an ordering vector in [0001] and an anti-ordering vector in $(11\bar{2}0)$ direction. The ordering is driven by a strong repulsive interaction of neighboring Sb atoms along the $(11\bar{2}0)$-oriented atomic chains on the $m$ plane sidewall facets.

In order to fabricate two-dimensional single WZ structure III-V semiconductor monolayers, [0001] oriented WZ InAs NW segments, were overgrown in-situ by InAs$_{1-x}$Sb$_x$ at 410°C using the Au-droplet assisted gas source molecular beam epitaxy. After growth, a thin capping layer of As was deposited onto the nanowires’ surfaces to prevent oxidation during transport through air into the analysis system. There, the nanowires were heated under UHV conditions to remove the capping layer. After cleaning, the NWs were cleaved-off and transferred to a freshly cleaned Si(111) surface, to investigate their sidewalls in-situ by STM.$^{[2,9]}$

Figure 1(a) illustrates schematically the structure of the investigated NWs, with the WZ structure InAs core in blue and the axially grown InAs$_{1-x}$Sb$_x$ ZB structure segment in red. As illustrated below, during ax-
of the InAs core. Second, the surface exhibits a high density of atomically sized protrusions (see magnification in Fig. 1(c)), arising from the presence of substitutional Sb atoms on As lattice positions. The contrast arises primarily from the outward relaxation induced by the Sb atoms with larger atomic radii as compared to the As atoms of the host lattice, in analogy to Sb atoms in ZB structure (110) III-V surfaces.

At closer look, all protrusions exhibit identical symmetries and intensities, centered exactly on top of a surface As sublattice position. Considering the crystal symmetry, substitutional Sb atoms in different layers in and below the surface exhibit different symmetries of the Sb contrast on the surface (Fig. 1(d)). Protrusions centered on top of surface As sublattice sites are compatible with Sb atoms incorporated on As sites in the surface layer, as schematically illustrated in the atomic model for the case labeled SbI in Fig. 1(d). In contrast, a Sb atom in the second layer (SbII) is centered in between of four surface As atoms, which can be anticipate to relax outward due to the larger Sb atom (as indicated by the larger size of the slightly lighter blue balls representing the neighboring As atoms in Fig. 1(d)). A count of the Sb protrusions yield concentrations ranging between 6.3 – 9.8% for our NWs.

The above discovery of a single monolayer shell raises the question, whether (short range) chemical ordering can emerge and how it might look like in such two-dimensional WZ structure InAs1−xSbx monolayer sheets. For identifying a possible short range order, we mapped the spatial distribution of the group V elements in the InAs1−xSbx monolayer shell with atomic resolution. This chemical map is then used for an in-depth analysis of the spatial correlation of the Sb distribution. Figure 2(a) illustrates the two-dimensional pair-correlation function (PCF) c(x, y) for Sb atoms. Values close to one represent statistically expected Sb pair distributions without interactions. Hence the data shows that at larger distances no interactions are present. However, for nearest neighbor Sb pairs along the c direction the PCF exhibits values much smaller than 1, indicating a low probability of occurrence (for quantitative comparability see Fig. 2(b)). In contrast, nearest neighbor Sb pairs in c direction, i.e. across the atomic chains, exhibit PCF values larger than 1, indicating more frequent occurrences than statistically expected. The results emphasize that a short range ordering, defined by an ordering vector (2z) in [0001] and an anti-ordering vector (2zAO) in [1210] di-
For a further understanding of the underlying physical mechanisms driving the ordering, we derive the so-called mean force potential \( W(x, y) = -kT \ln[c(x, y)] \) \cite{27}, which is a first approximation of the pair interaction potential in the low density limit. \( k \) is the Boltzmann constant and \( T \) the freeze-in temperature, below which no diffusion or site changes within the InAs\(_{1-x}\)Sbx monolayer occur anymore. An upper limit for \( T \) is assumed to be 623 K, the temperature used for evaporation of the As cap. The resulting mean force potential, shown in Figs. 2(c) and (d), reveal a strong Sb-Sb repulsion with a repulsive energy of about 35 meV in \( a \) direction and a Sb-Sb attraction with an attractive interaction energy of about 12 meV in \( c \) direction. Hence, the energy difference between both configurations is approximately \( 47 \pm 11 \) meV.

This value is compared to density function theory computations of the total energy difference of an InAs(10\(\overline{1}0\)) surface layer that exhibits nearest neighbor Sb-Sb pairs in either \( c \) or \( a \) direction. The calculations were carried out using the Vienna ab-initio simulation program (version 5.2.11). \cite{28} All electron projector augmented wave (PAW) pseudopotentials were taken \cite{29} using the Ceperley and Alder approximation, \cite{30} parameterized by Perdew and Zunger for exchange and correlation energies (i.e. the Sb, As, In, H.25 and H.75 PAW potentials). \cite{9} We used a kinetic energy cutoff of 312.5 eV. The calculations were performed for (10\(\overline{1}0\)) oriented supercells with different sizes of \( 4 \times 4, 4 \times 5 \), and \( 5 \times 5 \) and thicknesses of 8 as well as 10 monolayers (H terminated on back side). The supercells were optimized until forces on atoms were smaller than 0.9 meV/Å. The calculations yield similar total energies for all supercells. The average energy difference for Sb-Sb nearest neighbors in \( c \) and \( a \) direction is found to be 24 meV, with pairs in \( c \) direction being lower in energy. Note, nearest neighbor Sb-Sb pairs along the zigzag chains on the cubic ZB (110) surfaces exhibit a very similar bonding structure and indeed a very similar interaction energy of 33 meV was calculated and experimentally verified. \cite{9}

The calculated and experimental energy differences for pairs in \( c \) and \( a \) direction are in the same order, especially the energy for pairs in \( c \) direction is lower than for those in \( a \) direction. This can be understood by considering the the incorporation of Sb atoms with larger atomic radii in the WZ crystal structure: In \( c \) direction the bonding structure is less rigid and nearest neighbor Sb atoms on the surface are in fact second nearest neighbor group V atoms (neglecting the interbonded group III atoms), since only every second (0001) plane is visible in filled (or empty) state STM images (see schematic in Fig. 1(d)). In contrast, for Sb pairs along the \( a \) direction the Sb atoms are much closer since they are nearest neighbor group V atoms separated only by one group III atom. This enhances strain for such Sb pairs, in analogy with similarly bonded Sb pairs along the zigzag chain on ZB structure (110) surfaces. \cite{9}

At this stage we compare quantitatively the measured and calculated energy differences between the Sb pairs in \( c \) and \( a \) direction. The experimental value is somewhat larger than the calculated one. However, the experimental determination suffers from the uncertainty of the freeze-in temperature and from many body effect in the pair interactions. First, the freeze-in temperature is an upper limit and thus lower in reality. Hence, the measured energy difference is likely overestimated and can be anticipated to be lower by roughly 10 to 15 meV. Second, at the given Sb concentrations and interaction ranges, many body effects in Sb-Sb interaction can be anticipated to be present, since the ratio between concentration and interaction length is in analogy to that of dopant atoms in GaAs, where many body effects were shown to affect pair correlation functions. \cite{31} Hence, the mean field potential cannot be assumed to be equal to the pair interaction energy anymore and thus many body effects affect the measured energy difference. Although a
InAs nanowires by direct atomically resolved elemental mapping using scanning tunneling microscopy. The ordering is driven by strong Sb-Sb repulsions along the atomic chains in a direction, giving rise to ordering in non-polar directions, deviating from ordering in bulk WZ structure material. This highlights the importance of spatial confinement for the formation of novel chemical ordering structures in III-V semiconductor alloys.

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Figure 3. Perspective view of a ball model of the two-dimensional idealized (long range) chemical ordering of the monolayer shell on the InAs sidewall facet, based on the ordering and antordering vectors extracted from pair correlation function. (a) Side and (b) top view with surface unit cell (gray shaded area) and ordering as well as anti ordering vectors shown as green and red arrows, respectively.

quantitative agreement of the measured and calculated pair interaction energies is difficult to reach due to the mentioned effects, both, the experimental and theoretical results lead to the same physical conclusion, i.e. the presence of a novel short range order in a two-dimensional WZ structure InAs$_{1-x}$Sb$_x$ monolayer shell.

Now we turn to the idealized (long range) ordering structure. On basis of the ordering and antordering vectors, extracted from the pair correlation function, we derive the idealized chemical ordering structure of the monolayer shell on the InAs sidewall facet. The resulting ordering structure is shown in side and topview in Fig.3(a) and (b), respectively. The ordering vectors are indicated and give rise to lines of only Sb atoms along the $\langle 0001 \rangle$ direction alternating in $\langle 12\overline{1}0 \rangle$ direction with pure As atom lines. This idealized structure is named Iuliacumite, derived from the Latin name of the city of Jülich. [Lillite is already preoccupied as name. :-) ]

Finally, this raises the question why the sidewall overgrowth by WZ structure InAs$_{1-x}$Sb$_x$ is limited to only one monolayer. This limitation implies that nuclei of islands on top of the single monolayer shell are unstable and hence Sb atoms prefer incorporation in the axially grown upper ZB structure InAs$_{1-x}$Sbx section. This is conceivable, since InAs$_{1-x}$Sb$_x$ prefers the ZB structure even at very small Sb concentrations. However, overgrowth on sidewalls facets of WZ structure InAs NWs forces InAs$_{1-x}$Sb$_x$ to crystallize in the WZ structure of the InAs core. In addition, an inward diffusion of Sb is energetically prohibited$^{[22]}$ hence stabilizing the single monolayer InAs$_{1-x}$Sb$_x$ WZ structure shell.

In conclusion, we identified a novel short range chemical ordering type in single monolayer InAs$_{1-x}$Sb$_x$ wurtzite structure shells on $m$ plane sidewall facets of