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Composition profiling of InAs quantum dots and wetting layers by atom probe tomography and cross-sectional scanning tunneling microscopy

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This study compares cross-sectional scanning tunneling microscopy and atom probe tomography. We use epitaxially grown self-assembled InAs quantum dots (QDs) in GaAs as an exemplary material with which to compare these two nanostructural analysis techniques. We studied the composition of the wetting layer and the QDs, and performed quantitative comparisons of the indium concentration profiles measured by each method. We show that computational models of the wetting layer and the QDs, based on experimental data, are consistent with both analytical approaches. This establishes a link between the two techniques and shows their complimentary behavior, an advantage which we exploit in order to highlight unique features of the examined QD material.

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I. INTRODUCTION

Accurate and high-resolution structural imaging and compositional analysis techniques have been the key driving technological force behind recent advances in nanotechnology and nanoanalysis. Today, a multitude of analysis tools are employed by scientists and engineers studying nanostructures, each offering its own specific capabilities and advantages. Because each technique can supply different data, when a complex material or structure is being considered it is common that several methods will be used together in order to build a more complete and accurate understanding of the subject.

Of the various methods in use, there are two techniques that are of particular interest thanks to their unparalleled ability to provide atomic-level imaging. The first technique, cross-sectional scanning tunneling microscopy (XSTM), one of the family of scanning probe microscopy techniques, is well suited for studying semiconductor materials, particularly in the III-V (Ref. 1) and II-VI (Ref. 2) arena. The key strength of XSTM is that it can directly visualize the atomic structure of a material, allowing detailed structural analysis.3,4 However, analysis of nanostructures can be difficult due to the lack of contrast between atomic species, the outward relaxation of the surface, and the two-dimensional nature of the technique. Despite this, the great precision of the measurements makes it feasible to infer details that cannot otherwise be directly visualized. For example, statistical analysis and finite element (FE) calculations can be used to create three-dimensional (3D) models of the atomic structure.5,6 However, this FE approach, in which the outward relaxation of a strained surface is calculated, has an inherent limitation that it is not injective; any measured dataset can be simulated by a range of different input models.7

The second technique under consideration is atom probe tomography (APT). This is the latest evolution of the venerable field-emission microscope,8 whereby a field ion microscope is combined with a spatially resolved time-of-flight mass spectroscope, creating a device known as a three-dimensional atom probe (3DAP).9 In a 3DAP a needle-shaped specimen of material is placed under a large pulsed voltage causing single ions to be emitted from the specimen’s apex and accelerated toward a detector screen. The data collected after each voltage pulse can be combined to form a three-dimensional tomographic image of the specimen. This technique works best for materials with a high conductivity, where subatomic precision can be achieved, and so traditionally there has been little use for the technique outside of metallic systems.10,11 However, the re-emergence of the laser-based 3DAP has greatly expanded the range of materials that can be analyzed.12 Where semiconductor materials were previously out of bounds, now the additional thermal excitation from a pulsed laser makes performing APT on these materials feasible, although with a reduced analytical quality.13 Consequently, pioneering work to study semiconductor nanostructures, such as quantum dots (QDs), has started to be undertaken.14

In terms of capabilities, APT seems to complement XSTM very well. Where XSTM can image only 2D cross sections, APT provides a 3D tomographic reconstruction, and where XSTM has limited capabilities to distinguish between chemical species, the mass-spectral analysis offers the ability not only to distinguish between different elements but also different isotopes. Naturally, these capabilities do not come without a price. The length scales of the volume that can be measured by the 3DAP are typically of the order of tens of nm laterally and hundreds of nm vertically, which is far more restrictive than the many μm that can be imaged with XSTM. Furthermore, the spatial precision is not as great as with XSTM, and not all the emitted ions can be detected. Even in ideal conditions detection efficiency is less than 60%.15 Finally, a particular weakness of APT is that the images of the sample must be reconstructed, a process requiring many assumptions about factors such as apex shape, radius, evaporation conditions, and so forth.16,17 The result is that, while the 3DAP provides a very unique dataset, the reliability and spatial accuracy is inherently inferior to the direct measurements of XSTM.

II. EXPERIMENTAL DETAILS

The studied material is comprised of $\text{In}_x\text{Ga}_{1-x}\text{As}$ self-assembled QD layers in GaAs, grown on a (100) GaAs...
FIG. 1. (Color) Comparative view of typical data obtained with the two techniques under consideration. The five \((\text{In}_{x}\text{Ga}_{1-x})\text{As}\) layers are clearly visible by both methods. All the scale bars are 50 nm. (a) 190-nm × 165-nm topographic XSTM image taken at \(V = -3\) V, \(I = 30\) pA. Several QDs can be distinguished as bright features in the WLs. The thin line parallel to the QD layers is an adatom that is being pushed by the tip in the slow scan direction. (b) Atom map showing only the In atoms, marked as indigo pixels. The In atoms between the WLs are background noise and account for approximately 0.01–0.06% of the detected atoms. Red isosurfaces show the \(x = 25\%\) In fraction, marking the location of QDs. The top inset shows the reconstruction, now also with 20% of the detected Ga atoms (yellow pixels), overlaid on a scanning electron microscope (SEM) image of the FIB fabricated specimen prior to evaporation. The bottom inset shows the remains of the specimen after evaporation.

From atomic force microscopy (AFM) measurements on uncapped material grown under the same conditions, the areal density of the QDs was determined to be \(\approx 3.0 \times 10^{10}\) cm\(^{-2}\). Low-temperature (5 K) macro-photoluminescence measurements show that the QDs emit at an energy of 1.229 eV with a full width at half maximum (FWHM) of 59 meV.

The APT measurements were performed using an LEAP 3000X Si instrument, operating in laser mode. The laser...
wavelength was green (532 nm) with a pulse frequency of 0.5 MHz (2 \mu s) and a target evaporation rate was set to be 0.2%. The flight length between specimen and detector was 90.0 mm and the typical time of flight of an evaporated ion was 0.6\textendash1.6 \mu s, depending on the species. The evaporation rate refers to the probability that, for a given laser pulse, an event will be recorded by the detector. In this experiment therefore we aimed to achieve a single detection event every 500 pulses, that is, on average there would be 1 ms between each ion strike. The evaporation field is dynamically altered during the measurement in order to maintain this target. In general, the field will gradually increase over the course of the experiment as the radius of the specimen tip becomes larger as the specimen is evaporated down. Because the position sensitive detector in the 3DAP is not able to accurately distinguish multiple impacts it is important that the evaporation rate is low to maintain a high ratio of single-ion events.

The laser pulse energy was less than 0.01 nJ; such a low power was used in order to ensure good data quality.\textsuperscript{19} The sample stage temperature was set to 50 K. The needle-shaped specimen was fabricated via standard focused ion-beam (FIB) techniques using a dual-beam FEI Nova 200 NanoLab system.\textsuperscript{20,21} Prior to FIB sharpening a protective 100\textendash150-nm Ni layer was sputtered on top of the GaAs sample. Inside the FIB system, a further 100\textendash150-nm Pt layer was deposited on the Ni using the ion-beam deposition capabilities of the FIB.

All XSTM measurements were performed at room temperature under UHV ($p < 6 \times 10^{-11}$ mbar) conditions with an Omicron STM-1, TS2 Scanner. Electrochemically etched tungsten tips were used. The STM was operated in constant current mode on \textit{in situ} cleaved (110) surfaces. To ensure that the electronic contribution to the apparent height, as measured by XSTM, is minimized, a high negative bias voltage ($\approx -3$ V) was applied during all measurements resulting in a purely topographic signal\textsuperscript{22} allowing the identification of individual In atoms.\textsuperscript{23}

The strain relaxation, induced by the lattice mismatch between InAs and GaAs, of the WL and the QDs was modeled with the finite-element (FE) method. The FE calculations were performed using the MEMS module of COMSOL Multiphysics. To calculate the strain relaxation of the WL and the QDs a 2D and a 3D model was used, respectively.

### III. RESULTS

A comparative view of typical data obtained with the two techniques of XSTM and APT is shown in Fig. 1. From this figure, it immediately becomes apparent that the two techniques are of a different nature; where the XSTM measurement is restricted to a cleavage plane and thus yields information that is 2D in nature, the APT measurement provides a fully 3D dataset. Even without further analysis, the WLs and QDs can already be distinguished from Figs. 1(a) and 1(b). In the following sections, a detailed comparison is given of the WLs and the QDs, as measured by XSTM and APT.

#### A. Wetting layer

An atomically resolved 52-nm \times 19-nm local mean equalization filtered current XSTM image of a typical part of the WL is shown in Fig. 2(a). The WL is found to start abruptly (within one bilayer), followed by a decay of the In concentration in the direction of growth. From previous work it is known that the decay of the In concentration can be modeled by an exponentially decaying function.\textsuperscript{24} The function $a \exp(-z/b)$, with $z = 0$ the start of the WL and positive $z$ representing the growth direction, was used as an input for the FE modeling that was employed to calculate the outward relaxation profile [blue line, Fig. 2(b)], matches the outward relaxation profile as measured by XSTM.
[red line, Fig. 2(b)], the decay of the In concentration can be determined. We find $a = 0.188 \pm 0.002$ and $b = 1.95 \pm 0.05$.

The In concentration in the direction of growth can also be determined from the APT data. This is found by sampling the composition of 20-nm-diameter cylinders, aligned parallel to the $z$ axis. A small bin length was chosen in order not to lose detail of the sharp concentration changes. The thin diameter was chosen so the cylinder could be positioned not to coincide with any QDs and yet remain close to the core of the specimen, where the reconstruction is most accurate. The thin diameter also helps to reduce the effects of curvature in the reconstruction and misalignment of the cylinder to the normal axis, both of which could result in loss of detail in the sharp onset of the WLs. In order to ensure that there were no systematic errors in the reconstructions, volumes from different layers and different specimens were compared. To improve the signal-to-noise ratio, the concentrations of these different sampling volumes were averaged together; these averaged data are shown in Fig. 2(c) (top).

From the XSTM measurements we unambiguously determined that the onset of the WL is abrupt (within one bilayer). This is not the case in the APT data, where the In fraction rises from $x = 2\%$ to $15\%$ over a distance of $\Delta z \approx 0.7$ nm. Despite the measures taken to ensure a good interface, we consider this artifact to arise from the effect of averaging several volumes, which also do not individually have completely sharp interfaces, causing some error in determination of the start of the WL. However, since we know that the WL starts abruptly, we can perform a deconvolution of the APT profile in order to compensate for this. The deconvolution function was determined by fitting a Gaussian function to the onset of the WL in the APT data; see Fig. 2(c) (top). The FWHM of this Gaussian function is 0.8 nm and thus can be considered as the instrument profile. This value compares favorably with the resolution obtained from laser-assisted evaporation of Si isotope superlattices. An alternative technique, known as $z$-density correction, can also be used to improve interface sharpness in the depth direction.

The deconvoluted APT data are shown in Fig. 2(c) (bottom). The onset of the WL is now found to occur within $\approx 0.35$ nm, which is well within one bilayer (0.565 nm), and the peak indium fraction $x$ has risen by $\approx 2\%$, in agreement with the XSTM data. An exponentially decaying function in the same form as the input for the FE calculations was used to fit the deconvolved APT data. We find $a = 0.189 \pm 0.004$ and $b = 1.95 \pm 0.05$.

Comparing the two techniques we find an excellent match of the In segregation profiles. However, to arrive at this result a deconvolution of the APT data was necessary. In case no deconvolution function is known and the $z$-density correction is not applicable, the APT’s ability to image sharp interfaces is limited without better data reconstruction techniques.

**B. Quantum dots**

Having shown, with the aid of FE calculations, that the two techniques give comparable results for the WLs, we shall now consider the QDs. A total of 55 QDs, 43 by XSTM and 12 by APT, were characterized.

In Fig. 3, the height and width of all observed QDs is plotted. The height of the QDs was determined from the XSTM data by counting the number of bilayers in the current images, resulting in the discrete nature of the plotted data. In APT the height of the QDs was determined from a 1D composition profile sampled by a 10-nm diameter cylinder going through the center of the QD. The width is determined from the cross-sectional contour map with a 1-nm projection taken through what is estimated to be the longest part of the dot. An example of the
position of the cross section and example measurements of major and minor axes are shown in the upper-left and upper-center panels of Fig. 3(b), respectively. The thickness of the highest QD as measured by XSTM is in agreement with the highest QD observed by APT (2.8 nm vs. 2.7 nm). At the lower end of the observed height distribution a discrepancy between the two techniques is observed; the thinnest QD found by XSTM is significantly lower than the thinnest QD observed by APT (1.1 nm vs. 1.5 nm). This can be explained by the 2D nature of the XSTM technique. Due to the arbitrary position of the cleavage plane we have no a priori way of knowing how a QD is cleaved. If a QD is cleaved near its edge it will appear thinner. The same holds for the width of the QDs. This results in a wide range (7–22 nm) of observed QD widths and in the trend of increasing QD height with QD width in the XSTM data seen in Fig. 3(a).

In the fully three-dimensional technique of APT the height and width of the QDs can be determined with less ambiguity. It is therefore not surprising that the height/width distribution of the QDs as measured by APT is clustered in the top right of Fig. 3(a). The spread in the distribution reflects the nonuniformity of the growth process. By taking cross-sectional contour maps through noncentral positions of the QD, as shown in the upper-right panel of Fig. 3(b), we can emulate the effect of off-center cleavage. The yellow points in Fig. 3(a) show the width and height, measured as the contour map is moved toward the edge of the QD. In contrast to the blue points, where a 1D profile was used to determine the height, in this case it is measured from the $x = 25\%$ In concentration contour; this is not as accurate, but it gives more localized measurements.

It should also be noted here that there is a substantial error in the width of the QDs with both techniques. Due to the gradual transition of WL to QD it is hard to pinpoint where the QD starts. In an effort to increase the accuracy, a combination of current and topographic data was used to determine the width of the QDs with XSTM. In the case of the APT the WL profile is first subtracted from the data. However, with both techniques, an error of $\approx \pm 2$ nm remains in the width of the QDs. For reason of clarity these error bars are omitted from Fig. 3.

Figures 4(a) and 4(b) show the topographic and the local mean equalization filtered current XSTM image of a cleaved QD. The height and width of this QD was determined to be 2.8 and 18.0 nm, respectively. As mentioned above, it is normally not known how the cleavage plane intersects the QDs with the XSTM technique. However, the statistics of Fig. 3 show that this QD is one of the highest and widest that was observed. Therefore we can reasonably assume that this particular QD was cleaved through its center. Since not all the individual In atoms can be identified in the XSTM images, it is not possible to extract directly an In profile from Figs. 4(a) and 4(b), and an indirect route via APT and FE calculations has to be taken. With APT the In profile of the QDs can readily be obtained. A contour plot of the In concentration of a 1-nm-thick slice through the center of a QD measured by APT is shown in Fig. 4(c). The APT data were used to construct a model (d) which serves as an input for the FE calculations.
APT study has reported the peak In concentration in InAs QDs to be off-centered and toward the top of the QDs. This is not the case in our QDs where the peak In concentration is slightly inclined toward the top and laterally located in the center.

Conventionally, InAs/GaAs QDs are modeled by a disk or a truncated pyramid with increasing (from bottom to top) or inverted-triangular-shaped In profiles. Given the XSTM and APT data, such an approach cannot be followed with the current QDs. Figure 3(b) shows no strong evidence of faceting or a particular footprint for these QDs. As can be seen from the contour concentration maps, the QDs merge into the WL, making edges indistinct, and each has a unique irregularity. Therefore we consider it to be a reasonable first approximation to model these QDs as long thin spheroids with a circular footprint.

The size and shape of buried InAs/GaAs QDs are determined by a delicate interplay between driving and quenching of QD leveling during overgrowth. Depending on specifics of the growth conditions, the obtained QDs can be classified into two general types. The first class is characterized by a sharp interface between the QD and the surrounding matrix and is the result of a low growth temperature or a low growth rate. In the case of a low growth temperature the segregation of In out of the WL is strongly suppressed, resulting in a localized WL. In contrast to this, if the first class of QDs is formed by a low growth rate then a pronounced WL will be present. The second class of QDs is associated with a growth process involving a high growth temperature and a high growth rate. Under such growth conditions there is a strong In segregation during the overgrowth of the QDs. This results in a pronounced WL and softening of the interface of the QD with the surrounding matrix.

From Figs. 4(a) and 4(b) we observe that the interfaces of our studied QDs are not sharp and a pronounced WL is present. It is therefore clear that these QDs do not belong to the first class and are instead examples of the second class of QDs. We would like to note that it can be problematic to distinguish between these two classes in most XSTM and TEM studies of InAs/GaAs QDs, thus particular care must be taken when trying to model the QDs. To aid in the construction of a model of our QDs, two 1D In profiles were extracted from the APT data shown in Fig. 4(c): one through its center along the growth direction, shown in Fig. 5, and one laterally through the center, perpendicular to the growth direction (not shown). These data are fitted with analytical functions and scaled to match the width and height of the QD observed by XSTM depicted in Figs. 4(a) and 4(b). The analytical expressions were then extended in order to generate a fully 3D model of the QD. The result is shown in Fig. 4(d).

Besides the In profile in the core of the QD, two other regions proved crucial in modeling the QD: first, the exponential decay of the In concentration in the direction of growth above the QD. From the 1D In profile taken through the center of the QD, we determined that this amounts to 16% of the total number of In atoms present; see the marked area in Fig. 5. Note that in the XSTM images of Figs. 4(a) and 4(b), only a few individual In atoms are visible above the QD and an exponential decay cannot be distinguished.

Because the area above the QDs measured by XSTM is very limited a meaningful statistical analysis is not practicable. Consequently, the In atoms in this region are often overlooked. However, the APT data reveal that, together, these In atoms actually form a significant part of the indium profile and should thus be included in the model. The second region that is crucial in modeling our QDs is the gradual transition of the QD into the WL. The XSTM and APT images of Figs. 4(a)–4(c) show that the In concentration at the sides of the QD is more substantial than that of the WL profiled in the previous section. We choose to model this feature as a region of $x = 15\%$ In fraction which extends well beyond the QD and eventually merges with the WL [not shown in Fig. 4(d)].

In Fig. 6, the outward relaxation of the cleaved surface across the center and 7 nm off center of the QD, as measured by XSTM, is plotted together with the result of the FE calculations.

![FIG. 5. (Color) 1D In profile in the direction of growth through the center of the QD shown in Fig. 4(c). Analytical functions (solid lines) are fitted to the APT data. The In found above the QD in this profile is marked by the blue shaded area and amounts to 16% of the total amount.](attachment:fig5.png)

![FIG. 6. (Color) Outward relaxation of the cleaved surface of the QD shown in Figs. 4(a) and 4(b). The outward relaxation measured by XSTM through the center (red line) and 7 nm off center (yellow line) of the QD is compared with the results of the FE calculations (blue lines).](attachment:fig6.png)
calculations. A close match between the measured and the calculated outward relaxation is observed. Here it should be noted that different input models e.g., the models with a linear In profile mentioned above, can yield a similar match. However, such models would not resemble the APT data. From this we conclude that care is required when constructing QD models based on XSTM measurements alone. Recently, it has been shown\textsuperscript{7} that PL measurements in combination with extensive theoretical modeling and XSTM measurements had to be combined to yield a realistic QD model. However, the subtleties of the decaying In concentration above the QD and the extension of the QD into the WL might be overlooked in such an approach, and thus some ambiguity still remains. In our view, the abilities of APT in providing a 3D In profile are at the moment unique, and necessary to construct a valid 3D QD model.

IV. CONCLUSION

We have used XSTM and APT to image InAs/GaAs QD layers in a single sample. We have found there is good agreement between the segregation profiles of the WL obtained by both techniques. To arrive at this result it was necessary to deconvolve the APT data with an instrument profile. The height and length of the QDs determined by both techniques is also comparable, both in the case where cross sections are taken through the center of the QD and also where they are taken off-center. This highlights the versatility of APT whereby the 3D data can be processed in a variety of ways to show a variety of details. Exploiting this advantage, we have used the In profile measured by APT to make a 3D model of a typical dot. This model is in agreement with the outward relaxation measured by XSTM, and we therefore consider this to be a unique solution. This analysis method highlights some structural features of the QDs that were undetected or neglected in previous measurements. The juxtaposition of the two techniques shows the unique benefits and capabilities of each one. Thanks to their very different natures, by using XSTM and APT together we can analyze semiconductor nanostructures at a level of detail that has been previously unobtainable.

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