Rational synthesis of atomically thin quantum structures in nanowires based on nucleation processes

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Excitonic properties in quantum dot (QD) structure are essential properties for applications in quantum computing, cryptography, and photonics. Top-down fabrication and bottom-up growth by self-assembling for forming the QDs have shown their usefulness. These methods, however, still inherit issues in precision integrating the regimes with high reproducibility and positioning to realize the applications with on-demand quantum properties on Si platforms. Here, we report on a rational synthesis of embedding atomically thin InAs in nanowire materials on Si by selective-area regrowth. An extremely slow growth rate specified for the synthesis demonstrated to form smallest quantum structures reaching nuclear size, and provided good controllability for the excitonic states on Si platforms. The system exhibited sharp photoluminescence spectra originating from exciton and bi-exciton suggesting the carriers were confined inside the nuclei. The selective-area regrowth would open new approach to integrate the excitonic states with Si platforms as building-blocks for versatile quantum systems.

Semiconductor quantum dots (QDs) have attracted much attention as a light source using single and entangled photons in quantum cryptography¹,². The typical method of growing QDs is the self-assembled method using Stranski–Krasnov (S–K) mode due to lattice mismatch³. This growth method, however, has several challenges in controlling size, density, and positions and the S–K mode has been restricted in typical heterostructures such as In(Ga)As/GaAs systems. This limits the possibility of using optically active QDs on Si platforms for future Si-based optical circuits with versatile quantum system.

Recent progresses in epitaxial methods, such as vapor–liquid–solid method and selective-area epitaxy, have resulted in the growth of axial heterostructures in nanowires (NWs) regardless of lattice mismatch, i.e., various materials, such as quantum disks and QDs, could be embedded inside host NWs⁴–⁸. NW axial heterostructures have great potential for future quantum optical applications because flexible band-gap engineering is possible, and the diameter, height, and density of QDs can be controlled by modifying the host NW’s geometry. NW axial heterostructures have been fabricated in axial InP/InAs⁹, GaAs/GaP⁹, and GaAs/GaAsSb NWs¹⁰. These quantum structures embedded in NWs allowed single-electron transport⁹, resonant tunneling regime⁹, strong anti-bunching¹¹, and exciton spin memory¹¹. Moreover, tuning of optically active QDs’ charge state using gate voltage has been achieved using these NWs¹¹. The GaAs/InAs system as quantum light source, which operated at near-infrared wavelength region, are quite feasible for quantum system on-chip integration with Si based avalanche photo diodes inside Si microchips. These state-of-the-art phenomena induced by the quantum structures with GaAs/InAs axial heterostructures should be integrated on Si platforms as a future element in Si-CMOS compatible optical circuits using quantum cryptosystems.

In this regard, integration of QDs with Si and light-emitting diodes on GaAs/Si wafer-bonded substrates have been reported¹⁵,¹⁶. Embedding quantum structures into the host NW materials regardless of abnormal deposition on NW sidewalls is required for precise controlling and tuning of the quantum states on Si platforms. And the NW with nanoscale footprints would be expected to have strong confinement either or both in the axial and radial direction, which allows robustness and reproducibility of the excitonic properties. Thus, synthesis of an

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atomically thin quantum structure is essential for strong confinement. In addition, precise positioning of the structures is also important for fabricating on-demand light-source devices. Recent advances in selective-area epitaxy have enabled integration of vertical GaAs NWs and light sources on Si substrates\(^\text{17,18}\), and templated assisted selective epitaxy (TASE)\(^\text{19,20}\), which uses Si microstructures as templates, enables the integration of III-Vs nanomaterials on Si nanostructures such as electronic\(^\text{21}\) and optical\(^\text{22}\) devices on Si platforms. By using GaAs NW’s grown using selective-area epitaxy as templates for regrowth, atomically thin InAs layers can be embed inside these thin NWs. Therefore, we characterized vertical stacked GaAs/InAs/GaAs heterostructures in position-controlled GaAs NWs on Si by the selective-area regrowth.

**Results**

**Selective-area regrowth of atomically-thin InAs embedded GaAs NWs on Si substrates.** We demonstrated selective-area regrowth to embed four InAs layers inside vertical GaAs NWs on Si. This method prevents unintentional growth on the sidewalls of host NWs by covering the sidewalls with amorphous (SiO\(_2\)) films. We discuss the growth morphology of atomically-thin InAs layers by the selective-area regrowth. The grown InAs layers exhibited sharp photoluminescence (PL), indicating the formation of InAs quantum structures (disks or dots). And excitation power dependence on the PL intensity revealed that the InAs atomically thin layer via nucleation of two-dimensional (2D) InAs islands resident exciton and bi-exciton.

Vertical GaAs NWs were grown on Si(111) by using selective-area epitaxy by changing the initial Si surface into a (111)B-polar surface\(^\text{17}\). The vertical GaAs NWs array was successfully integrated on the Si(111) surface by the selective-area epitaxy (Fig. 1a). The design of the GaAs/InAs/GaAs axial heterostructures consisted of four InAs layers, the growth cycle of each InAs was 2, 6, 10, and 20 cycles with 10-nm-thick GaAs barriers. The processes of selective-area regrowth were as follows. First, 100-nm-thick SiO\(_2\) was deposited on the vertical GaAs NWs, as shown in Fig. 1b. In the sputtering procedure, the sidewalls of the GaAs NWs were covered with 50-nm-thick SiO\(_2\). Next, reactive-ion-etching (RIE) was carried out to etch the top portion of the SiO\(_2\)-covered NWs (Fig. 1c). Then, the top part of the NWs were selectively etched using NH\(_4\)OH:H\(_2\)O\(_2\):H\(_2\)O = 5:1:1 solution. This etching process was used to form a SiO\(_2\) cylinder on the GaAs NWs, as illustrated in Fig. 1d. The etching solution selectively formed GaAs(111)A surface inside the SiO\(_2\) cylinder. Then, GaAs buffer layer was grown under the same growth condition of GaAs NW in order to recover the GaAs(111)B top surface. Finally, GaAs/InAs/GaAs vertical stacked layers were grown by pulsed growth with different InAs cycles, as shown in Fig. 1e. The purpose of the pulsed growth for the GaAs/InAs/GaAs layers was to elongate the surface diffusion length of the Ga and In atoms inside the SiO\(_2\) nanotube and avoid abnormal deposition outside the sidewalls of the SiO\(_2\).

**Photoluminescence of the atomically-thin InAs structures.** Figures 2a showed the growth results of the InAs atomically-thin layer embedded GaAs NWs on Si substrate. We designed InAs layers with four different numbers of cycle, as illustrated in Fig. 2b. A uniformly SiO\(_2\)-coated GaAs NW consisting of the four InAs layers, illustrated in Fig. 2b, was integrated on the Si substrates. The NW diameter was 100 nm, height was approximately 1 \(\mu\)m, and diameter fluctuation was \(\pm\)4 nm. There were no abnormal depositions on the SiO\(_2\) sidewalls, which indicated the source materials react on only the (111)B surface surrounded by the SiO\(_2\) tube. The micro (\(\mu\))-PL spectra under low excitation power in Fig. 2c showed sharp optical emissions around 1.30 eV at the excitation power of 0.92 kW/cm\(^2\). The number of sharp PL peaks corresponded to the number of InAs layers. The other PL peaks with broad full width at half maximum (FWHM) at around 1.30 eV were assumed to be deep level emission from the GaAs NWs.

The positions of the four sharp PL peaks were 1.342 (peak A), 1.334 (peak B), 1.312 (peak C), and 1.277 eV (peak D). These PL peak positions were almost constant at arbitral positions in the GaAs/InAs/GaAs NW array
Figure 2. Atomically thin InAs layers-embedded GaAs NWs on Si. (a) 45°-tilted SEM image showing the InAs-embedded GaAs NW array on Si. (b) Illustration of GaAs/InAs/GaAs multilayers grown using the selective-area regrowth. (c) μ-PL spectra for GaAs/InAs/GaAs NWs on Si measured at 4.2 K. (d) Peak deconvolution of the PL spectra using Lorentzian function. (e) Plot of transition energy for heavy-hole and light-holes as a function of InAs thickness for GaAs/InAs/GaAs strained QW on GaAs. (f) Plot of InAs thickness with variation in the pulse cycles.
(see in Supplementary Fig. S1). This means the size fluctuation of the InAs layer is significantly small in the regrowth process. We analyzed the PL spectra by using Lorentzian functions for the sharp PL and a Gaussian function for the broadened PL. The curve-fitting with Lorentzian function estimated FWHM of 630 μeV for peak A, 1.2 meV for peak B, 1.7 meV for peak C, and 980 μeV for peak D shown in Fig. 2d. These PL properties exhibited similar luminescence behavior to the other InAs QDs which was formed by S–K growth and selective-aera growth (see in Supplementary Table S1).

Figure 2e shows the calculated transition energy level in a strained GaAs/InAs single QW. We ascribed the InAs QW on GaAs(111)24 and excluded the quantum confinement effect in the radial direction. Assuming the sharp PL peaks in Fig. 2c originated from the InAs layers, the InAs thickness was linearly changed with increasing the InAs cycle (Fig. 2f). The growth rate of the InAs layers was estimated to be 0.1 Å/s, which was extremely slower than 1 monolayer [ML, 3.498 Å for InAs(111)]. This indicates that the InAs ML-growth proceed with very slow growth rate involving a step flow growth in the SiO2 tubes.

Discussions

Nucleation growth process in selective-area regrowth of the atomically thin InAs layers. We investigated the early stage of InAs SAG on GaAs(111)B surface to reveal the extremely slow rate in the InAs regrowth process. Figure 3a,b show the SAG of InAs on GaAs(111)B surface, the growth time of which was 5 min. Growth condition for the InAs was almost same as InAs layer (explained in method), but the gas supply was continuous flow condition without pulsed growth. Figure 3a illustrates the coalescence of triangular-shaped InAs islands on large openings with a opening diameter of 400 nm. On the other hand, hexagonal-shaped structures were formed on openings with a diameter of 100 nm (Fig. 3b). In case of conventional SAG, the nominal growth rate was approximately below 1 Å/s for InAs (and 10 nm/min for GaAs) which was estimated from that on SAG of InAs (GaAs) NWs on GaAs(111)B substrates. Although the growth in the InAs/GaAs lattice mismatch system proceeded with the S–K mode involving a wetting layer, the InAs on the GaAs(111)A surface exhibited formation of 2D stacking fault tetrahedron25. The InAs nucleation process of SAG on GaAs(111)B thought to follow the formation of the 2D islands.

Similar growth morphology was observed in the other heteroepitaxial system in such GaAs/Si (lattice mismatch is about 4.1%)17. Another investigation of STM reported that the formation of 2D InAs islands26. The 2D InAs islands have flat (111)B surfaces, and the initial stage of the InAs NW growth on GaAs(111)B was based on the coalescence of these InAs 2D islands because multinuclear growth and next stacking proceed before the InAs monolayer (ML) is fulfilled the whole surface in the openings. Note that a part of the initial GaAs(111)B surface thought to have As-trimer27 resulting in a slow nucleation rate due to the surface reconstruction. And the InAs crystallization gradually starts from the steps of the InAs 2D islands. Figure 3c shows the average height of InAs NWs as a function of growth time, indicating the height was sub-linearly increased with growth time due to the above growth process. The dashed lines are guide lines of 1 Å/s and 128 Å/s, respectively. In the early growth stage, the growth rate was much smaller than the 1 Å/s, which coincides to the growth rate estimated in Fig. 3b.

Figure 3d illustrates growth evolution of the atomically thin InAs layer. With increasing the number of pulse cycle for InAs regrowth, the first nuclei (2D InAs island) eventually form a complete ML through the step-flow wavefunction. When the InAs pulse cycle was 10, 2ML-thick with 2D InAs islands as illustrated in Fig. 3d(v) were observed in the vicinity of the top surface. The 2D InAs layer showed around 3–10-nm-width lamellar strain, which are never explained only by the rotation twin. The width of the lamellar strain field was covered with a section of the whole surface. This abnormal strain field thus suggested the formation of ML-thick 2D InAs islands, which was speculated in Fig. 3d. In case of InAs layer grown with 18 cycles [the yellow square (ii) in Fig. 3g], the similar lamellar strain estimated from the 3ML-thick InAs was observed. These images indicated that the second 2D InAs island was elongated on the 2ML-thick InAs layer via step-flow growth, which was ascribed in Fig. 3d(v).
Optical transition process for the atomically thin InAs.

Figure 4a shows the excitation power dependence of each PL band. These PL bands showed double-splitting with increasing excitation power density. This splitting relates to exciton/biexciton or ground-state/higher state levels in InAs layers. Figure 4b shows the integral PL intensity for each PL band as a function of excitation power density. The blue circles are the lower positions of PL spectra (A1 ~ D1). The red circles are the higher peaks (A2 ~ D2). In the PL spectra (A and B), the PL intensities in A1 and B1 monotonically increased with increasing excitation power density. This excitation-power dependence indicates the optical transition originating from free exciton emission. The PL intensities for A2 and B2, however, quadratically increased with increasing the excitation power density, meaning biexciton emission was involved. These behaviors of the exciton and biexciton emissions were similarly observed in other NWs such as InAsP quantum dots embedded in InP NWs and InAs QDs grown on GaAs NWs.

The energy level of biexciton is typically lower than that of exciton because of the positive binding energy for biexciton resulting from attractive Coulomb interaction. However, the biexciton in the regrown InAs layer had negative binding energy, exhibiting a higher energy state than that of exciton. This means that the Coulomb
interaction of the two localized excitons is repulsive due to the narrow confinement space. The negative binding energy was 10 meV in InAs QDs \(^3\) and the energy depends on the shapes of the QDs \(^3\). The energy difference between the exciton and biexciton for bands A and B at an excitation power of 0.96 kW/cm\(^2\) was −6 and −3 meV.

Figure 4. Optical properties for the atomically thin InAs layers in GaAs NWs. (a) Excitation-power dependence of μ-PL spectra of GaAs/InAs/GaAs NWs on Si. (b) Integrated PL intensity for bands A, B, C, and D as a function of excitation-power density. (c) Temperature dependence of μ-PL spectra. Excitation power density was 0.96 kW/cm\(^2\).
reaches the nucleation process in the selective-area regrowth, the growth technologies would open new scheme early growth stage of 2D InAs islands with slow growth rate. By utilizing the extremely slow growth rate that originated from exciton emission of atomically thin 2D InAs islands.

The selective-area regrowth involves specific InAs heterointerface.

radiative lifetime was shorter than those of PL band A, which was related with misfit dislocation of the GaAs/GaAs layers. The partial pressures of TMIn and AsH3 were 4.8 × 10^{-7} and 1.3 × 10^{-4} atm, respectively. The partial pressure of the GaAs was same as those of the GaAs NWs. The GaAs buffer layer was the same condition as the GaAs NWs. The InAs and GaAs were grown by pulsed growth. The growth temperature for GaAs/NWs multilayers were 560 °C. A cycle of the InAs and GaAs growth was carried out with Group III supply (TMIn or TMGa) for 1 s and AsH3 supply for 1 s with H2-intervals for 1 s.

Figure 4c shows the temperature dependence of the PL spectra. The narrow PL emissions remained at the same intensity between 4.2 and 40 K, then decreased with an activation energy of ~ 52 meV for A1 and 61 meV for A2, and consequently quenched at ~ 70 K. The activation energy was estimated using an Arrehnius equation. (The fitting results for bands A and D are shown in Supplementary Fig. S4.) The activation energy for band A corresponds to those of (111)-oriented InAs QDs. The rate parameter for the A1 and A2 were 5.9 × 10^5 and 1.75 × 10^6, respectively. The large activation energy for band A indicates efficient electron–hole capture and radiative recombination in the 2D InAs island. The smaller rate parameter for band A1 means the thermal emission transport from the 2D into underneath the InAs layer was efficient.

The activation energy for other PL bands, such as B, was estimated to be 17 meV for bands B1 and 12 meV for B2 B2. In the band B, the activation energy for B1 was larger than band B2, both energies remained at 10 meV. This indicated that almost all photo-generated carriers were thermally transport to underneath the InAs layers with increasing temperature. The activation energy for band D1 and D2 were estimated to be 13 and 32 meV, respectively, while the rate parameter was 50 and 380, respectively. These small rate parameters indicate that the radiative lifetime was shorter than those of PL band A, which was related with misfit dislocation of the GaAs/InAs heterointerface.

In conclusion, we have demonstrated selective-area regrowth of forming atomically-thin InAs layers in GaAs NWs integrated on Si platforms. The atomistically thin InAs layers inside GaAs NW have exhibited sharp PL peaks originated from exciton emission of atomically thin 2D InAs islands. The selective-area regrowth involves specific early growth stage of 2D InAs islands with slow growth rate. By utilizing the extremely slow growth rate that reaches the nucleation process in the selective-area regrowth, the growth technologies would open new scheme in integrating the quantum regimes with Si platforms as building-blocks for versatile quantum systems.

Methods
Selective-area growth of vertical GaAs NWs. After the Si(111) was degreased with organic solvents, 20 nm-thick SiO2 film was formed by thermal oxidation. Hexagonal-shaped openings arranged in a triangular lattice with a pitch of 0.4–1 μm were then formed on the amorphous films by using electron-beam (EB) lithography and wet chemical etching. The opening diameter d0 was 70 nm. Metalorganic vapour phase epitaxy (MOVPE) was carried out to grow NWs. The GaAs NWs were grown in a horizontal low-pressure MOVPE system. Trimethylgallium (TMGa) and arsine (AsH3) gas were used as material sources. After thermal cleaning at 900 °C in H2, we carried out AsH3 treatment at 400 °C and low-temperature GaAs were grown for 3 min at 400 °C. Then, the GaAs NWs were grown for 60 min at 760 °C. The partial pressures of TMGa, AsH3 were respectively 1.0 × 10^{-5} and 2.5 × 10^{-4} atm.

Formation of GaAs/InAs layers for selective-area regrowth. Trimethylindium (TMIn) and AsH3 were used as the materials for InAs growth. The partial pressures of TMIn and AsH3 were 4.8 × 10^{-7} and 1.3 × 10^{-4} atm, respectively. The partial pressure of the GaAs was same as those of the GaAs NWs. The GaAs buffer layer was the same condition as the GaAs NWs. The InAs and GaAs were grown by pulsed growth. The growth temperature for GaAs/InAs multilayers were 560 °C. A cycle of the InAs and GaAs growth was carried out with Group III supply (TMIn or TMGa) for 1 s and AsH3 supply for 1 s with H2-intervals for 1 s.

Microphotoluminescence measurement. Micro-PL (μ-PL) measurement was carried out at 4.2 K. The excitation light was He–Ne (632.8 nm) focused to a 2-μm spot using a 100 × objective lens. About ten NWs were inside the excitation laser spot at the same time. The PL spectra were taken with a nitrogen-cooled charge-coupled device detector.

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Author contributions

K.T. designed the experiments, made the nanowires by MOVPE, measured all luminescence data and evaluated analyzed all of the data. T.F. and J.M planned and supervised the study. All authors discussed the results and commented on the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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