Growing a LaAlO$_3$/SrTiO$_3$ heterostructure on Ca$_2$Nb$_3$O$_{10}$ nanosheets

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The two-dimensional electron liquid which forms between the band insulators LaAlO$_3$ (LAO) and SrTiO$_3$ (STO) is a promising component for oxide electronics, but the requirement of using single crystal SrTiO$_3$ substrates for the growth limits its applications in terms of device fabrication. It is therefore important to find ways to deposit these materials on other substrates, preferably Si, or Si-based, in order to facilitate integration with existing technology. Interesting candidates are micron-sized nanosheets of Ca$_2$Nb$_3$O$_{10}$ which can be used as seed layers for perovskite materials on any substrate. We have used low-energy electron microscopy (LEEM) with in-situ pulsed laser deposition to study the subsequent growth of STO and LAO on such flakes which were deposited on Si. We can follow the morphology and crystallinity of the layers during growth, as well as fingerprint their electronic properties with angle resolved reflected electron spectroscopy. We find that STO layers, deposited on the nanosheets, can be made crystalline and flat; that LAO can be grown in a layer-by-layer fashion; and that the full heterostructure shows the signature of the formation of a conducting interface.

Transition metal oxide (TMO) perovskites form an interesting group of materials with a large variety of physical properties, among which the occurrence of superconductivity, ferromagnetism, and of ferroelectricity$^1$. Moreover, stacking layers of different such oxides allows for new properties to be made on design. A well known example is the formation of the two-dimensional electron system (2DES) between the band insulators SrTiO$_3$ (STO) and LaAlO$_3$ (LAO)$^2,3$. Typically, these interfaces are fabricated by growing a layer of one material (LAO) on a single crystal substrate of the other (STO). The formation of the conducting interface requires the STO crystal or film to be terminated by its TiO$_2$ layer, on which the AlO$_2$ layer of the LAO can grow. The polar discontinuity between the TiO$_2$ layer with its net zero charge and the AlO$_2$ layer with a charge of $-e$ (e the electron charge), together with oxygen defects forming during the deposition of the LAO layer, are two important ingredients in the charge transfer to the interface and the formation of the 2DES$^4,5$. This is the reason that STO substrates are almost exclusively used in researching the 2DES properties. Since only a thin layer of SrTiO$_3$ is needed to create the 2DES, integration with other materials could profit from the ability to use different substrates. Growth of STO on Si has been demonstrated$^6$, and conditions were found which allow epitaxial growth, but it would still be useful to have as few constraints as possible for the choice of substrate. Here we develop that approach and use inorganic nanosheets. Such sheets can be obtained through exfoliation of materials with a lattice closely matching the lattice parameter $a_0$ of SrTiO$_3$ ($a_0 = 0.3905$ nm). In our case that is Ca$_2$Nb$_3$O$_{10}$ (CNO; $a_0 = 0.386$ nm), on which the growth of epitaxial SrTiO$_3$, SrRuO$_3$ and BiFeO$_3$ has already been demonstrated$^7$. For STO in particular, it was found that STO grows unstrained, due to the flexibility of the CNO nanosheet. This approach is very flexible, as nanosheets can be deposited on virtually any substrate$^7-10$, rendering design freedom in tailoring device properties. The exfoliated nanosheets have a thickness of only 1.5 nm to 4 nm$^{10-13}$, which means that they can also be used in experiments which make use of a backgate.

To study the growth of LAO/STO on CNO we use low-energy electron microscopy (LEEM) with in-situ pulsed laser deposition (PLD). This combination allows us to follow details of the growth in real time but also to investigate the electronic structure of the material as we grow it. We recently reported on a LEEM study of the growth of LAO on STO with various terminations$^{14}$, and we showed that (i) we can observe layer-by-layer growth

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through the oscillations in the spot width and intensity of the specular beam in the diffraction pattern as function of time; (ii) we can image the layer morphology, in particular surface atomic steps; and (iii) most importantly, we can obtain an electronic fingerprint from the LAO surface during growth, which allows us to determine whether this layer is giving rise to a conducting or to an insulating interface. The fingerprint is based on angle-resolved reflected-electron spectroscopy (ARRES), a technique which allows to map the empty electron bands of the layer at energies above the vacuum energy. By comparing the results obtained here on nanosheets with our earlier study of LaAlO₃/SrTiO₃, we come to the conclusion that we can grow flat and crystalline heterostructures, and that they show the fingerprints of a conducting LaAlO₃/SrTiO₃ interface.

Results

Growth of the STO template layer. We discuss four different samples. On samples A we study growth during STO deposition; sample B is used to characterise and improve the surface of the deposited STO layer before depositing LAO; samples C is used to grow 5 unit cells of LAO on top of the STO layer; and sample D is a reference samples where again 5 unit cells of LAO is grown on a TiO₂-terminated STO substrate. Figure 1a shows a bright field LEEM image of Ca₂Nb₃O₁₀ nanosheets deposited on Si (sample A). Individual nanosheets can clearly be recognized. The square surface net expected for Ca₂Nb₃O₁₀ is visible in the LEED pattern (Fig. 1b) taken from the same area as Fig. 1a. Note that the substrate does not contribute LEED spots because of the native oxide on the Si surface. Besides the four principal spots originating from the big flake covering the bottom half of the image, many other rotated patterns are visible originating from the flakes in the top half of the image. This indicates that the nanosheets are randomly oriented on the Si substrate. The rotations of individual flakes can be determined by recording μ-LEED patterns. For example, Fig. 1c,d show that the sheets indicated with green (top) and red (bottom) circles in the LEEM image in Fig. 1a have different orientations. The diffraction spots are not very sharp, due to imperfect crystallinity or absorbates at the surface.

On the sheets we grew STO by PLD at a growth temperature of 660 °C. Higher temperatures showed degradation of the diffraction pattern. In order to follow the evolution of the surface during growth, we recorded IV-curves betwenn 0 eV and 25 eV after every 5–10 laser pulses. Figure 2 shows a color representation of the evolution of the intensity and full-width-half-maximum (FWHM) of the specular diffraction spot (the (0,0) spot) taken from those IV-curves. During growth we observe a clear change of the intensity (Fig. 2a) and therefore the FWHM becomes energy dependent. The FWHM is given in percentage of the size of the Brillouin zone.
a change in the electronic fingerprint of the surface, which first of all indicates that deposition is taking place. The IV-curves converge to a constant spectrum after around 300 pulses which consists of maxima around 3 eV, 8 eV and 15 eV. As we will discuss below, these are spectra typical for SrTiO$_3$. The FWHM (Fig. 2b) is a sensitive measure of the crystallinity of the surface, while disorder yields broadening of the (0,0) spot over all energies. Surface roughness due to the presence of unit-cell size step edges, only influences the FWHM at energies where the incoming and reflected electrons destructively interfere along the out-of-plane axis of the crystal$^{17}$. During growth, the diffraction spots become sharper, quantified by a global reduction of the FWHM (Fig. 2b) and the FWHM becomes energy-dependent. This leads us to conclude that the crystallinity improves with time, but the surface is still rough.

Importantly, neither the intensity nor the FWHM showed growth oscillations, while we observed such oscillations during the growth of STO and LAO on STO$^{14,18}$. For the growth on nanosheets both intensity and FWHM are constant, and the FWHM is large. It is not clear whether this difference is caused by a small coherence length in the material or a non layer-by-layer growth mode with a constant rough surface, but it precludes us from an accurate measurement of the deposition rate. Comparing the growth conditions and amount of pulses to earlier measurements, we estimate a film thickness close to 3.3 nm or 8.5 unit cells. This means the IV-curves do not change anymore after about 7 unit cells (300 pulses).

Figure 3a shows that the deposited layer (presumably STO) grows epitaxially on the nanosheets despite this roughness, from a comparison of a section of the (0,0) spot for the original nanosheets (blue, dashed) after annealing, and after growth of SrTiO$_3$ (green, solid) on sample A taken at $E_a = 17$ eV. The intensity was normalized, meaning the dashed curve was multiplied by roughly 8 to be visible on the same level as that of the green curve. (b) Bright field image after growth of 9 u.c. STO at $E_a = 29.6$ eV with locations of μ–LEED measurements d and e indicated in red (left) and green (right). (c) LEED pattern of the area shown in (b). (d), (e) μ–LEED patterns recorded in the areas indicated in (b). The LEED/μ–LEED images in (b–d) were taken at $E_a = 17$ eV. Figure taken from$^{16}$. 

![Figure 3](https://www.nature.com/scientificreports/)

**Figure 3.** (a) Normalized section of the specular diffraction spot on pure nanosheets (blue, dashed) after annealing, and after growth of SrTiO$_3$ (green, solid) on sample A taken at $E_a = 17$ eV. The intensity was normalized, meaning the dashed curve was multiplied by roughly 8 to be visible on the same level as that of the green curve. (b) Bright field image after growth of 9 u.c. STO at $E_a = 29.6$ eV with locations of μ–LEED measurements d and e indicated in red (left) and green (right). (c) LEED pattern of the area shown in (b). (d), (e) μ–LEED patterns recorded in the areas indicated in (b). The LEED/μ–LEED images in (b–d) were taken at $E_a = 17$ eV. Figure taken from$^{16}$. 

![Diagram](https://www.nature.com/scientificreports/)

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Fig. 3d,e show single orientations for every STO/nanosheet heterostructure, demonstrating that the SrTiO3 follows the crystal orientation of the underlying nanosheets.

The LEEM-IV spectra can be used to further characterize the deposited layer. Figure 4a shows IV-curves of the (0,0) spot, taken on sample B (STO thickness of about 5.7 nm, i.e. 600 PLD pulses). The lowest (blue) curve was taken after deposition, to be compared to a reference curve taken on a TiO2-terminated STO substrate (green; 2nd curve from the top) and a curve taken on a SrO-terminated STO substrate (magenta; top curve). The reference curve was taken from ref. 14, while the SrO-termination was obtained by the growth of an SrO bilayer on a TiO2-terminated substrate. The signature of the as-deposited surface clearly shows more resemblance to the SrO-terminated surface, with an intensity minimum around 12 eV and a broad maximum around 15 eV. In a next step, we heated the samples at constant rate from 660 °C to 770 °C during 30 min, and holding the temperature constant for another 30 min at a constant oxygen pressure of 5 × 10⁻⁹ mbar (equal to the pressure used during PLD).

The annealing changes the surface electronic properties and also further improves the surface quality. Figure 4a shows the IV-curve of the (0,0) spot after annealing (red curve, 2nd from bottom). From the change, and in particular from the appearance of a minimum around 8 eV, we conclude that the surface changes from a dominantly SrO-termination to a TiO2-termination. The evolution of the IV curves as function of time is shown in more detail in Fig. 4b where it can be seen that there are clearly discernable changes around 20 minutes, that the 8 V minimum has developed around 30 minutes, and that no further changes occur during the next 30 minutes. Figure 4c shows the corresponding change of the FWHM of the specular beam. The FWHM clearly decreases in the first 10–20 minutes, while the energy dependence disappears, indicating that the surface flattened. Clearly, the change to a TiO2-termination, the increase in crystallinity, and the flattening of the surface go hand in hand.

Growth of LAO. Next, we use these improved surfaces as growth template for thin films of LaAlO3. In Fig. 5a we compare the IV-curves before (blue, bottom) and after (red, middle) growth on sample C (STO thickness of about 5.5 nm, i.e. 588 PLD pulses) with a reference curve of 5 unit cells LaAlO3 grown under the same conditions on an STO-substrate (sample D) (green, top). The IV-curves after growing LAO on the STO/nanosheet sample and after growing on the TiO2-terminated STO substrate are very similar, confirming that similar LAO films were grown in both cases. Moreover, and even more importantly, we observed clear oscillations in both the intensity and the FWHM of the specular beam during the deposition, demonstrating epitaxial growth in layer-by-layer mode. This is analogous to using Reflection High Energy Electron Diffraction (RHEED) in standard PLD growth experiments. The full evolution of the intensity and FWHM maps during growth is shown for sample C (Fig. 5b,c) and for the reference sample D (Fig. 5e,f). The intensity oscillations are particularly clear around Eₘ₀ = 16 eV, indicating that both samples grow in the same layer-by-layer fashion. The value of 16 eV is the same as found in earlier experiments, as expected, since it is determined by the energy (or wavelength) where impinging around step edges destructively interfere. As a guide to the eye, vertical dotted lines indicate the full unit cells, showing that 5 unit cells were grown. Growth oscillations are even more clear from the changes in the FWHM at 16 eV, which are plotted for the nanosheet sample in Fig. 5d. After growth, the shape of the nanosheets is still clearly visible. Figure 6 shows a LEEM (a) and an AFM image (b) taken on sample C. Line profiles taken on a single flake (Fig. 6c) show a surface roughness of less than a unit cell (Ra value), pointing to an atomically flat surface.

The question remains whether the LAO/STO heterostructure has the desired electronic properties. We studied the LEEM-IV signature of conducting interfaces before and we use those results here. The IV-curves for the LAO/STO/nanosheet sample (Fig. 5a) can be characterized as showing a peak around 9 eV, a dip around 13 eV,
and another (lower) peak at 17.5 eV. This is the characteristic signature for the conducting LAO/STO interface; for non-conducting interfaces, we find a much broader and higher peak around 18 eV and the suppression of the dip. A more robust distinction can be made by measuring the full angular dependence of the (reflected) intensity using the ARRES technique. In ARRES we measure electron reflectivity maps not only as a function of landing energy \( E \) but also of the in-plane wave vector \( k_\parallel \) of the electrons. The plot can be represented in the usual Brillouin zone terminology, which in this case spans the high-symmetry points \( \Gamma \), \( X \) and \( M \) of the square surface net. The reflected intensity is determined by the unoccupied states above the vacuum level. Here, unoccupied bands are presented as intensity minima while band gaps are maxima, because the electrons are reflected with high probability if no states exist in the material at that \( E \) and \( k_\parallel \).

Figure 5. (a) IV-curves before (blue, bottom) and after (red, middle) growth of LaAlO\(_3\) on sample C as well as a reference curve of 5 unit cells LaAlO\(_3\) on a SrTiO\(_3\) substrate on sample D (green, top). The curves are shifted for clarity. (b) Intensity map versus time and energy of the specular \((0, 0)\) diffraction spot on sample C. (c) the same map on reference sample D. (d) FWHM oscillations versus number of laser pulses (time) at \( E_\parallel \) = 16 eV, on sample C. This energy is close to the out-of-phase conditions where maximal surface sensitivity is reached. (e) FWHM map for all energies on sample C and (f) FWHM map on sample D. Figure taken from.

Figure 6. (a) Bright field LEEM image at \( E_\parallel \) = 16 eV and (b) AFM image of the LaAlO\(_3\)/SrTiO\(_3\)/Ca\(_2\)Nb\(_3\)O\(_{10}\) heterostructure (sample C). (c) Line profiles of the relative height difference on the nanosheets; profiles from top to bottom correspond in (b) to white bars from left to right. The dark area in the center of the LEEM image is due to an inhomogeneity in channel plate of the detector. Figure taken from.

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SrTiO3/nanosheets heterostructure (sample C) with the reference sample of LAO on a TiO2-terminated STO-substrate (sample D) and maps of an LaAlO3/SrTiO3 sample with a conducting interface (sample S1-C from ref. 14), and a non-conducting interface (sample S2-I from ref. 14), respectively. Both maps were taken after deposition of 4 unit cells in a deposition run which continued to 8 unit cells. It should be noted that the data for the nanosheet sample were acquired by taking bright-field real-space images and selecting a single flake in the post processing15, in contrast to the data of ref. 14 where a 20 μm2 area was averaged by using LEED data.

The salient features for a heterostructure with a conducting interface are a strong peak at the Γ-point around 9 eV, a weaker peak around 18 eV, and a strong minimum in between; plus high reflected intensity at the X-point for energies between 13 eV and 19 eV. This can be seen in both conducting samples in Fig. 7b,c. The intensity distributions are somewhat different, probably because the first is a 5 u.c. sample and the second a 4 u.c. sample. Around the Γ-point, the sample grown on the nanosheets (Fig. 7a) looks similar to these, and rather different from the insulating sample of Fig. 7d, which is characterized by an intensity peak at the Γ-point around 9 eV, and a roughly equally strong and also very broad peak (in k-space) around 18 eV, but little intensity at the X-point. Around the X-point, the difference between the nanosheet sample and the insulating sample are actually obvious, which is best seen by imagining a line between Γ and X at around 17 eV. The conducting samples and the nanosheet sample show a peak-dip-peak intensity variation, while the insulating sample just shows a diminishing intensity. On the whole, the X-point differences seem the most important but both from Γ-point and X-point considerations the nanosheet sample shows stronger similarities with the conducting sample S2-C (Fig. 7c).

**Discussion and Conclusions**

The first examples of perovskite growth on Ca2Nb3O10 nanosheets in the literature already showed they are promising candidates for transfer of TMO devices to literally any substrate. Here we have shown how low-energy electron microscopy studies can help to find the growth conditions and annealing steps required to build TMO devices on Ca2Nb3O10 nanosheets. In depositing an STO layer directly on top of the nanosheets, we did not observe intensity oscillations in the specular beam as expected for layer-by-layer growth. This is at least partly due to the growth conditions, more specifically the growth temperature. For the first STO layer we could not obtain a crystalline film at temperatures above 660 °C and degradation of the diffraction pattern was observed before the growth started. In the literature however, films are grown at 700 °C7,10. The main difference between the
conditions in refs. 7,19 and those reported here is the oxygen background pressure. From this we infer that nanosheets degrade in low oxygen pressures for temperatures above 660 °C. This could be caused by the reduction of oxygen in the nanosheet. On the other hand, the crystallinity of the STO layer clearly improved during growth of the first 7 unit cells and this number seems to be required for a well defined surface with SrTiO3-substrate-like properties.

The grown STO film showed a crystalline but rough surface, as can be concluded from the quite broad specular spot. This could be improved by annealing, which again suggests that the growth temperature was too low for ideal growth. The as-grown surface showed an IV-curves characteristic for an SrO -terminated STO sample, while after annealing at 770 °C the surface showed the IV-curve of TiO2-terminated STO. This suggests that the laser fluence was slightly too low to grow stoichiometric STO. It is well known that the laser fluence influences the Sr/Ti cation ratio. Working with a fluence below the one yielding a 1:1 ratio will result in a film with a (slight) Sr excess20. Growing in this regime of excess Sr can actually lead to a termination conversion when starting on a TiO2-terminated surface, as reported by Bauemer et al.21. Still, that annealing at 770 °C changes this is somewhat unexpected since the SrO-terminated reference curve was measured at that same temperature. However, the top layer of the SrO-terminated reference sample was a double layer of SrO forming a stable rock salt structure, and this may be different for the sample we grew here. We surmise that the SrO we observe on the grown SrTiO3 is due to the rough surface and low growth pressure, and unstable at high temperatures. Annealing reorders the surface to a well defined TiO2-termination.

The roughness, incidentally, also appears to preclude the occurrence of a surface reconstruction. When Ti-deficient growth is started on a flat STO surface, SrO islands form on which a 2x2 reconstruction is found22. This surface with half-unit-cell islands on top constitutes a mixed-terminated surface. We do not observe such a surface in Fig. 3b. The only statement we can make is that the surface shows the signature of an SrO termination.

While the nanosheets themselves were unstable at temperatures above 660 °C, no degradation is found during the annealing of the SrTiO3 film on nanosheets at 770 °C. Apparently the STO layer protects the nanosheets from oxygen reduction. However, the temperature dependence is still fragile and temperatures above 770 °C are not recommended at these low pressures. Once a well-defined surface was obtained, layer-by-layer growth of LaAlO3 could be achieved. In the behavior of the intensity oscillations during growth we did not find any difference between the growth of LAO on our STO film on nanosheets and a TiO2-terminated STO substrate. The final ARRES map is very similar to ARRES maps found on conducting LaAlO3/SrTiO3 heterostructures as found in ref. 14. From a close comparison with the measurements performed on samples with conducting and non-conducting interfaces we conclude that the sample grown on the nanosheets shows the characteristics of a conducting interface. This implies not only a TiO2-terminated STO layer, but also the correct LAO stoichiometry.

Lastly we stress the opportunity of electronic gating of TMO devices on nanosheets. We infer a minimum SrTiO3 buffer layer thickness of only ≈ 7 unit cells. For bottom gating of devices this would mean a gate dielectric of only 7 unit cells plus the thickness of the nanosheets, which was reported to be between 1.5 nm and 4 nm16–19. A gate dielectric of only 7 nm would provide significant possibilities for bottom gating.

**Summary**

We have used a Low Energy Electron Microscope with built-in pulsed-laser-deposition capability to study the feasibility of growing an SrTiO3/LaAlO3 bilayer with a two-dimensional electron system (2DES) at its interface on lattice-matching nanosheets of Ca2Nb3O10, which in turn were deposited on the native oxide on top of a Si substrate. Apart from electron diffraction as a characterization tool, we show how to use the intensity-versus-energy characteristics to obtain more information on the electronic structure of the layers we deposit, and we use Angle-resolved reflected-electron spectroscopy (ARRES) to study the electronic structure of the completed sample. We find conditions leading to a TiO2-terminated STO surface, we demonstrate that we can use intensity oscillations during growth in order to count the number of LAO unit cells, and we infer from ARRES maps that the surface of the sample shows an electronic structure similar to what was found for samples with a conducting interface. The versatility of the nanosheet approach should therefore allow local fabrication of 2DES systems with their own desired geometry.

**Methods**

Ca2Nb3O10 nanosheets were synthesized and deposited by Langmuir-Blodgett (LB) deposition on Si (001) substrates cut from single crystal wafers with native oxide layers as described elsewhere. The resulting (unilamellar) layer is about 1 to 2 nm thick. The samples were transferred into the ESCHER LEEM system23–26 with in-situ pulsed laser deposition as described in ref. 14. PLD growth and LEEM imaging in this system are performed alternately, since the high voltage between objective lens and sample has to be turned off during the sequence of laser pulses. Before starting the growth, the samples were annealed at 570 °C (measured with a pyrometer using an emissivity of 0.7) for at least half an hour in an oxygen background pressure of 5 × 10−5 mbar, in order to remove any contaminants.

For the deposition of SrTiO3 onto the nanosheets, the temperature was raised to 660 °C. PLD was performed with a laser fluence of 1.4 J/cm2 and a 1 Hz repetition rate in 5 × 10−5 mbar oxygen16. This may result in slightly off-stoichiometric growth of STO16, but in our microscope it is not possible to increase the pressure and further optimize the growth conditions. Following the deposition, the sample was annealed at 770 °C for at least half an hour in the same oxygen pressure, in order to improve the surface flatness. The sample was then cooled down and moved to the transfer chamber (2 × 10−8 mbar) while the STO target was replaced with an LAO target. After a pre-ablation step of the LAO target, the sample was transferred back to the measurement position. It was heated to 750 °C at the same oxygen pressure of 5 × 10−5 mbar. LaAlO3 was then deposited at a laser fluence of 2.0 J/cm2 and 1 Hz repetition rate. In ref. 14 it was shown that these parameters result in a conducting LAO/STO interface. Three samples were made with a specific number of laser pulses, namely 360 (sample A), 600 (sample B) and 580
The thicknesses we estimate from these pulse numbers, for reference purposes, are 3.3 nm (A), 5.7 nm (B) and 5.5 nm (C). As was already mentioned, we estimate thicknesses from comparing the growth conditions and amount of pulses to earlier measurements. In this way we find a film thickness close to 3.3 nm or 8.5 unit cells for sample A (360 pulses).

The nanosheet starting material, the growth process and the final heterostructures were investigated by various techniques available in the LEEM. In particular real-space imaging was used to identify nanosheets, μ-LEED to study crystallinity and roughness, and LEEM-IV spectra (the intensity of reflected electrons as a function of their landing energy \( E_\text{a} \))\(^{27}\) to identify various states of the growing surface. We also used angle-resolved reflected-electron spectroscopy (ARRES)\(^{15}\), which extends the concept of IV-curves by recording electron reflectivity maps as a function of landing energy and in-plane wave vector \( k_x \) of the electrons. Such ARRES maps are correlated with the unoccupied band structure of the surface layer and are therefore well suited to probe the surface stoichiometry of the LaAlO\(_3\)-layer. This implicitly indicates whether or not a 2-DEG is formed at the LaAlO\(_3)/\text{SrTiO}_3 interface as was shown before\(^{14}\).

One sample was imaged by Atomic Force Microscopy (AFM), using a standard commercial microscope (Bruker) in tapping mode.

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Author contributions
A.v.d.T., J.ten.E., G.K., S.v.d.M. and J.A. contributed to the conception and design of the experiment. H.Y., Z.L., J.ten.E., G.K., G.R. and M.H. were involved in the preparation and characterization of the nanolayers. A.v.d.T., M.H. and J.J. performed the PLD growth, the LEEM measurements and the data analysis. A.v.d.T., J.J., S.v.d.M. and J.A. wrote the manuscript, with assistance of the other authors.

Competing interests
The authors declare no competing interests.

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