Influence of Sr deficiency on structural and electrical properties of SrTiO₃ thin films grown by metal–organic vapor phase epitaxy

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Homoepitaxial growth of SrTiO₃ thin films on 0.5 wt% niobium doped SrTiO₃ (100) substrates with high structural perfection was developed using liquid-delivery spin metal–organic vapor phase epitaxy (MOVPE). Exploiting the advantage of adjusting the partial pressures of the individual constituents independently, we tuned the Sr/Ti ratio of the gas phase for realizing, stoichiometric, as well as Sr deficient layers. Quantitative energy dispersive X-ray spectroscopy in a scanning transmission electron microscope confirm Sr deficiency of up to 20% in nominally off-stoichiometrically grown films. Our MOVPE process allows to grow such layers in phase pure state and without extended defect formation. Indications for oxygen deficiency could not be identified. Sr deficient layers exhibit an increased permittivity of $\varepsilon_r = 202$ and a larger vertical lattice parameter. Current–voltage characteristics (IVCs) of metal–oxide–semiconductor (Pt/SrTiO₃/SrTiO₃:Nb) structures reveal that Sr deficient SrTiO₃ films show an intrinsic resistive switching with on–off ratios of three orders of magnitude at RT and seven orders of magnitude at 10 K. There is strong evidence that a large deviation from stoichiometry pronounces the resistive switching behavior. IVCs conducted at 10 K indicate a defect-based mechanism instead of mass transport by ion diffusion. This is supported by in-situ STEM investigations that show filaments to form at significant higher voltages than those were resistive switching is observed in our samples.

Strontium-titanate (SrTiO₃) has widely been investigated due to its interesting physical properties such as high permittivity at room temperature in bulk crystals, ceramics, and thin films²⁻³. While unstrained and undoped SrTiO₃ is a centrosymmetric material⁴ and paraelectric at room temperature, it can be transferred into a ferroelectric state by the application of strain⁵. By doping with niobium, lanthanum, or samarium, SrTiO₃ becomes an n-type semiconductor, which may even show superconductivity at very low temperatures as described by Ahadi et al.⁶ Especially in recent years, SrTiO₃ thin films have gained an increasing interest due to its resistive switching behavior, rendering it a potential candidate for resistive random access memory (ReRAM) applications⁷. However, the underlying physical origin of this effect is still controversial to date. Different models based on Schottky barriers⁸ and charge trapping effects⁹ were discussed to explain the bipolar resistive switching effect, yet the most commonly accepted models for the origin of resistive switching are oxygen migration models⁷,¹⁰–¹⁴. By application of a high bias pulse, conductive filaments of oxygen vacancies are formed and ruptured by diffusion processes, which leads to a low resistive state (LRS) and high resistive state (HRS), respectively. The impact of cation deficiency¹⁵,¹⁶ on soft-forming processes was investigated on thin films grown by pulsed laser deposition (PLD)¹⁷,¹⁸.

Growth of SrTiO₃ thin films by different techniques has often been reported in literature. The most frequently used method is pulsed laser deposition (PLD), but also (hybrid) molecular beam epitaxy (MBE)¹⁹,²⁰, sol–gel²¹, and metal–organic chemical vapor deposition (MOCVD)²²–²⁶ techniques have been utilized. While in the PLD process film composition and structure are determined by an interplay of different growth parameters, like laser fluence, geometry, background pressure and target composition, MBE and MOCVD offer the possibility
Here, the (Sr/Ti)liq ratio in the source materials was varied from 2.0 to 3.6. By controlling the Sr and Ti gas flows, the corresponding partial pressures—and particularly the ratio of the partial pressures—are precisely adjusted in corresponding complex oxides and hence are formed at lower Ts. When Ts is increased to 750 °C, the energy shows the corresponding HRXRD patterns of the thin films with the film reflection emerging as small shoulders.

In this paper, we demonstrate the appropriate growth regime for deposition of phase-pure SrTiO3 epitaxial thin films by MOCVD, which we will denote as metal–organic vapor phase epitaxy (MOVPE) in the following.

As a result of these investigations, we were able to realize coherent, homoepitaxial growth of phase pure SrTiO3 thin films on 0.5 wt% Nb doped SrTiO3 substrates. By independent control of Sr and Ti fluxes into the reaction chamber, we have precisely adjusted the Sr/Ti ratio in the gas phase and thus in the films, which allows us to systematically study the impact of Sr/Ti ratio on the structural and (di)electrical properties. Thereby, we found an increasing dielectric permittivity and a pronounced intrinsic resistive switching behavior at high Sr deficiency.

We reveal that the underlying point defects in the film are related to deliberately introduced Sr deficiency. Further, we report on intrinsic resistive switching behavior even at low temperatures of 10 K, where ionic diffusion processes should be strongly suppressed. Hence, resistive switching in non-stoichiometric films (strong Sr deficiency of 20%) shows an atypical evolution compared to reported resistive switching devices based on SrTiO3 materials, providing evidence of defect-assisted transition instead of a mechanism based on mass transport.

**Results and discussion**

Section A is focusing on the growth, stoichiometry and characterization of the defects present in the thin films, whereas section B is dealing with the impact of Sr deficiency on the permittivity and resistive switching. In Section C, *in-situ* STEM investigations are presented.

**Growth and stoichiometry control of SrTiO3 by MOVPE.** In order to determine the appropriate substrate growth temperature (Ts) for achieving single-phase SrTiO3 growth, depositions were performed from 550 to 750 °C. The thin films were grown on DyScO3 (110) substrates using a (Sr/Ti)liq = 1 with an absolute Sr concentration of cSr(tmhd)2 = 0.025 mol/l. DyScO3 was chosen here as substrate to distinguish film and substrate peaks in high-resolution X-ray diffraction (HRXRD). Figure 1a shows 2θ/ω scans of samples grown under various substrate temperatures Ts.

For growth temperatures below 675 °C, additional Bragg reflection peaks occur at 2θ of about 27°, 32°, 38°, and 43°, which are attributed to the single oxide contributions TiO2 (103), SrO (111), TiO2 (203), and SrO (104), respectively. Additionally, a SrCO3 (111) phase (2θ ≈ 26°) is formed at 650 °C. From 675 to 725 °C, single-phase SrTiO3 with (100) surface orientation was observed, whereas at a temperatures of 750 °C additional Ruddlesden–Popper (RP) phases like Sr2TiO4 and a TiO2 phase are present. The vertical lattice parameter of the SrTiO3 thin films occurs in the vicinity of the DyScO3 peak and thus at a lower 2θ angle compared to an unstrained, stoichiometric SrTiO3. This is attributed to an off-stoichiometry in the films, which will be discussed in more detail below. This temperature dependence of phase formation agrees with literature data and is reasoned by the standard formation enthalpies. Single oxides in general have lower formation enthalpies than their corresponding complex oxides and hence are formed at lower Ts. When Ts is increased to 750 °C, the energy provided is sufficient to form RP phases, which are known to have even higher standard formation enthalpies than SrTiO3. Thus, in order to obtain single-phase SrTiO3 without the formation of single oxides, carbonates, and RP phases, Ts should be adjusted in the range of 675 °C to 725 °C.

Next, we grew a series of samples on 0.5 wt% Nb-doped SrTiO3 (100) substrates using a fixed Ts of 710 °C. Here, the (Sr/Ti)liq ratio in the source materials was varied from 2.0 to 3.6. By controlling the Sr and Ti gas flows, the corresponding partial pressures—and particularly the ratio of the partial pressures—are precisely adjusted in the gas phase and consequently, the cation composition in the layer can be modified in a targeted way. Figure 1b shows the corresponding HRXRD patterns of the thin films with the film reflection emerging as small shoulders at the low angle side of the substrate peak. As achieved for the series grown on DyScO3, no parasitic phases like SrO, SrCO3, TiO2, SrTiO3, SrSi2Ti2O9, SrSi3TiO11 (n = 1, 2,...) appear for this growth temperature, demonstrating phase pure (100) SrTiO3 epitaxial films for all applied (Sr/Ti)liq ratios.

In Fig. 1c, a higher angular resolution of the HRXRD pattern around the (200) substrate peak is presented. The (Sr/Ti)liq ratios were below 3.2, the film peak occurs at lower 2θ angles as compared to the SrTiO3 substrate peak. Furthermore, for (Sr/Ti)liq > 2.6, occurrence of thickness fringes can clearly be observed, which indicates a high crystalline quality of the film including smooth interfaces and surfaces. However, the thickness fringes in the HRXRD patterns are overlaid with irregular oscillations for almost all films. As we will discuss later in conjunction with the STEM investigation, these oscillations are related to a fluctuating defect distribution along the growth axis and hamper the fitting of the HRXRD pattern, which assumes a single homogenous
layer. Rather, a stack of at least two films each with a different vertical lattice parameter is required to match measurement and simulation. For detailed insights into the HRXRD pattern simulations, see section HRXRD simulation in the Supplemental.

The evaluation of all HRXRD data indicates that the film peak is approaching the position of SrTiO₃ bulk substrate with increasing (Sr/Ti)liq ratio up to 3.6. From the angular difference of the Bragg reflection of the film and the substrate, the vertical lattice parameters \(d_\perp\) of all samples with \(2.0 \leq (Sr/Ti)_{liq} \leq 3.6\) are calculated and plotted in Fig. 1d. The plot reveals a continuous increase of the vertical lattice parameters \(d_\perp\) with decreasing (Sr/Ti)liq ratios. We attribute the enlarged lattice parameters of the films for (Sr/Ti)liq < 3.6 to an off-stoichiometry and a resulting compressive in-plane lattice strain in the films. Only for (Sr/Ti)liq = 3.6, the film lattice parameter matches the bulk lattice parameter of \(d_{SrTiO_3} = 3.905 \text{ Å}\), suggesting stoichiometric film composition and a strain-less film. The correlation between the increased vertical lattice parameter in our MOVPE thin film and the deviation from stoichiometric composition qualitatively agrees with the results from other deposition techniques. However, the observed deviations of the vertical lattice parameter in our MOVPE films from the value for unstrained, stoichiometric SrTiO₃ is much larger than for SrTiO₃ thin films grown by PLD or MBE methods as summarized by Brooks et al. indicating that we can achieve much higher off-stoichiometric conditions without occurrence of secondary phases. This is attributed to the fact that MOVPE is operated close to thermodynamic equilibrium: by the application of suitable substrate temperatures the formation of secondary phases is suppressed. This possibility to stabilize huge deviations from stoichiometric composition gives MOVPE an unique characteristic in comparison to all the other deposition methods.

In order to verify the epitaxial relation between film and substrate, we performed reciprocal space maps (RSM) for a non-stoichiometric film with (Sr/Ti)liq = 2.6 in the vicinity of the asymmetric (204) Bragg reflection of bulk SrTiO₃ (see Fig. 1e). The appearance of substrate and film peaks at the same in-plane reciprocal lattice parameter indicates pseudomorphic epitaxial film growth on the SrTiO₃ substrate.

To rule out that an oxygen deficiency is responsible for the systematic change of the lattice parameter in the nominally Sr deficient layers in the as-grown-state, we performed in-situ 2θ/ω HRXRD scans while annealing the samples in pure oxygen atmosphere up to 950 °C. HRXRD patterns were recorded for a sample grown at (Sr/Ti)liq = 3.0 at 30 °C, 200 °C, 500 °C and from 500 to 950 °C in 25 K intervals with one hour dwell time at each stage, as shown in Fig. 1f. For selected temperatures. After completing the described annealing cycle, another 2θ/ω HRXRD scan was again recorded at 30 °C. Neither the film peak position nor the shape or distance of the
A film with a (Sr/Ti)\text{liq} ratio of 3.6 shows near stoichiometry. STEM-EDX measurements on an off-stoichiometric film grown with a (Sr/Ti)\text{liq} ratio of 3.2. Figure 3a exhibits a contrast similar to an anti-site domain boundary and is discussed in detail in the Supplemental section. cloudy contrast. In comparison, films with a (Sr/Ti)\text{liq} ratio of 3.6 only show a faint contrast with respect to the substrate making the film-substrate interface indistinguishable in some regions. Under these imaging conditions, the intensity of the atomic column reflects the mean atomic number (Z-contrast). Therefore, we conclude that the spreading of the values determined in this way is ±1.2%, which corresponds to the measurement error. In particular, we notice a composition variation along the growth direction. This was also reflected in the different contributions to the HRXRD patterns, such as the smearing of the film peak and the superposition of several interfering patterns (Details see Supplemental). Notably, the STEM-HAADF contrast varies across the epitaxial layers of the films in Fig. 2, indicating a spatially inhomogeneous distribution of the underlying defects. In particular, we observe a stacking fault in the STEM-HAADF images, which is locally changing the channeling conditions and thus the corresponding Ti EDX signal. Nevertheless, if there is a lower Ti signal in the layer, its contribution is negligibly small. Towards the surface, the Sr content increases, which essentially results in an unintentional sub-layer structure. We have cross-checked such substantial Sr deficiencies by means of off-axis STEM-HAADF, yielding a comparable outcome (see Supplemental).

For a microscopic study of the structural properties and the stoichiometry, we investigated nominally Sr deficient layers grown with a (Sr/Ti)\text{liq} ratio of 2.6, 3.2, and the stoichiometric layer with a (Sr/Ti)\text{liq} ratio of 3.6 by TEM. The corresponding high-resolution scanning transmission electron microscopy (STEM) images are displayed in Fig. 2a–c, respectively. For the sake of comparability, the HAADF intensities in the region of the SrTiO₃ substrate were normalized to unity. All STEM-HAADF images confirm the formation of single-crystalline and coherently grown SrTiO₃ films without the presence of foreign phases, in agreement with the HRXRD observations. Furthermore, no extended defects or misfit dislocations could be observed by STEM. For the given HAADF imaging conditions (camera length of 196 mm), layers with a (Sr/Ti)\text{liq} ratio of 2.6 and 3.2 generally appear considerably darker as compared to the SrTiO₃ substrate with a cloudy contrast. In comparison, films with a (Sr/Ti)\text{liq} ratio of 3.6 only show a faint contrast with respect to the substrate making the film-substrate interface indistinguishable in some regions. Under these imaging conditions, the intensity of the atomic column reflects the mean atomic number (Z-contrast). Therefore, we conclude that films with a (Sr/Ti)\text{liq} ratio of 2.6 and 3.2 exhibit a reduced mean atomic number as compared to the substrate, which is in-line with a Sr deficiency in the layer as intended. In contrast, the faint contrast to the substrate of the film with a (Sr/Ti)\text{liq} ratio of 3.6 shows near stoichiometry.

To further corroborate the Sr deficiency in the layer, we carried out additional STEM energy dispersive X-ray (STEM-EDX) measurements on an off-stoichiometric film grown with a (Sr/Ti)\text{liq} ratio of 3.2. Figure 3a exhibits the color-coded EDX maps of the Sr, Ti, and O EDX signal, as well as the corresponding STEM low-angle annular dark field (HAADF) intensity (camera length 300 mm). While the Ti and O signals reveal no apparent change between the layer and the substrate region, the Sr signal in the MOVPE layer is substantially lower, which is particularly pronounced close to the interface. The STEM-LAADF map, which is sensitive under these conditions to diffuse scattering from strong atomic displacements along the beam direction, clearly correlates with the Sr EDX intensity. This implies that Sr vacancy and/or their related complexes are responsible for the observed lattice distortions. To quantify the Sr deficiency, Fig. 3b displays the horizontally averaged EDX signals of Sr (green line) and Ti (blue line) along the growth direction. We have quantified the precision of the EDX measurements by determining the 95% confidence interval of the Sr/Ti ratio values measured in the stoichiometric substrate. The spreading of the values determined in this way is ±1.2%, which corresponds to the measurement error. In the stoichiometric substrate region, which serves as reference, the Sr and Ti signals are practically identical. Within the epitaxial layer, the Ti as well as the Sr signal decrease in intensity, with the latter being much more pronounced. To quantify the decrease of Sr, Fig. 3b displays the ratio between the Sr and Ti signal (purple line), revealing Sr deficiency up to around 20% in the SrTiO₃ film close to the substrate interface. Regarding the slightly smaller Ti signal in the MOVPE layer, we believe that this effect is related to the strong atomic displacements, which are locally changing the channeling conditions and thus the corresponding Ti EDX signal. Nevertheless, if there is a lower Ti signal in the layer, its contribution is negligibly small. Towards the surface, the Sr content increases, which essentially results in an unintentional sub-layer structure. We have cross-checked such substantial Sr deficiencies by means of off-axis STEM-HAADF, yielding a comparable outcome (see Supplemental).
Electrical properties of Sr deficient layers. In this section, the influence of the Sr deficiency on the (di)electrical properties of the SrTiO$_3$ thin films is investigated. For this purpose, the MOVPE layers were measured in the metal–oxide–semiconductor (MOS) structure described in the methods section and schematically shown in the inset of Fig. 4. All layers were initially insulating for RT and 10 K measurements (> GΩ at a bias voltage of 0.1 V). The relative permittivity was measured at room temperature and showed no dependence on contact size. In Fig. 4, the permittivity is shown versus the vertical lattice parameter. The permittivity continuously increases in the range between $d_\perp = 3.907$ Å to $d_\perp = 3.956$ Å from 110 to 202 and then again decreases for samples with higher vertical lattice parameter. The film with $d_\perp = 3.956$ Å exhibits the highest permittivity, even higher than the film with stoichiometric composition ($d_\perp = 3.907$ Å). This evolution agrees qualitatively with literature data for films grown by PLD, MOCVD and sol–gel processes and is understood in terms of polar nanoregions and lattice distortions breaking the paraelectric cubic symmetry of stoichiometric SrTiO$_3$. Furthermore, it can be deduced that the permittivity cannot be regarded as a fixed material property since it critically depends on the amount of Sr deficiency.

For the electrical characterization, current–voltage characteristics (IVCs) were measured first at room temperature in the voltage range of $-4$ V $\leq V \leq 3$ V. The polarity is chosen as it is commonly done in literature with the back-contact on substrate side as common ground and the applied bias at the top-contact. These cycles were reproducible for several repetitions for both kind of samples.
Sr deficient samples with \((\text{Sr/Ti})_{\text{liq}} = 3.2\) showed the typical hysteresis behavior for intrinsic resistive switching (see Fig. 5a). Once the LRS or the HRS is enabled, it is possible to probe each state within the bias range of \(-0.1\ \text{V} \leq V \leq 0.1\ \text{V}\) without any change of resistivity (insets Fig. 5a). The on–off ratio defined as \(I_{\text{LRS}}/I_{\text{HRS}}\) at 0.01 V is about \(10^3\) for the off-stoichiometric sample.

In contrast, for the stoichiometric thin films with \((\text{Sr/Ti})_{\text{liq}} = 3.6\), stable resistive switching could not be observed in this applied voltage range (see Fig. 5b). The resistivity at read-out voltage of 0.1 V stays at a high level, similar to the initial resistivity of an as grown sample.

At bias voltages larger than 0.5 V a significant current increase can be observed. At such high field strengths of up to 1 MV/cm, injected charge carriers into the film play a role in electrical transport of the thin film, e.g. by Schottky\(^4\) or Poole–Frenkel emission\(^4\). A detailed analysis of the electrical transport is beyond the scope of this work and will be given elsewhere.

Next, current–voltage measurements of non-stoichiometric films with \((\text{Sr/Ti})_{\text{liq}} = 3.2\) were performed at low temperature using a closed-cycle refrigerator (\(T = 10\ \text{K}\), He background pressure = \(10^{-6}\ \text{mbar}\)) to demonstrate resistive switching. At low temperature, the switching voltage is more clearly visible and remains with about 1.5 V essentially unchanged compared to room temperature.

Another crucial observation is the soft forming process at low temperature. For this, the voltage range was gradually increased until intrinsic resistive switching is introduced. Within the first three sweeps (sweep #1 to #3 in inset of Fig. 6), the voltage is stepwise increased to the threshold voltage of \(-1\ \text{V}\). At this point, no resistive switching is observed. Only starting from sweep #4 (\(-2\ \text{V} \text{ to } 1.5\ \text{V}\)), partial resistive switching could be observed. By further increase of the voltage (sweep #5), the on–off ratio increases to about seven order of magnitude. After

![Figure 5. IV curves for samples with two different values of (a) \((\text{Sr/Ti})_{\text{liq}} = 3.2\), and (b) 3.6. The IV curves are measured in the range of \(-4\ V \text{ to } +3\ V\) showing two different states of resistivity. The upwards measurement in HRS is shown as red line, while the downwards measurement is shown as blue line. The probing of the state (HRS/LRS) is shown in each plot as inset.](https://www.nature.com/scientificreports/)

![Figure 6. Current voltage measurements at 10 K for SrTiO\(_3\) thin film with \((\text{Sr/Ti})_{\text{liq}} = 3.2\). The voltage range was gradually increased until resistive switching occurs (Sweep #1 to #6). The curves of sweep #1 to #3 are shown vertically shifted to each other by a factor of 10 in the inset. (Partial) switching occurs for \(V \text{ > } 1.5\ \text{V}\).](https://www.nature.com/scientificreports/)
reaching the hysteresis of sweep #5 (−4 V to 2 V) intrinsic resistive switching is stable for all following cycles, as illustrated by sweep #6. As shown, the hysteresis evolved without forming pulse and gradually without any abrupt change. Remarkably, the HRS for IVs displaying (partial) switching (#4 to #6) show a higher resistivity compared to the IV measurements below the threshold (#1 to #3). This is especially visible for negative bias. This observation is in strong contrast to the soft-forming process in Sr deficient resistive switching materials, as described by Lenser et al.13 and Stille et al.44. There, the overall conductivity of the sample increases since oxygen vacancies are incorporated during the first sweep, and hence implement additional charge carriers. In our particular case of Sr deficient MOVPE thin films, we observe the opposite, thereby providing evidence of an alternative mechanism for resistive switching, which is not based on ion diffusion. The contribution of ion diffusion can be neglected for our IV measurement conditions at 10 K. In addition, we can exclude intrinsic resistive switching based on antiphase boundaries as shown by Hensling et al.45 The existence of antiphase boundaries could not be verified by TEM. Further, due to our Ti-rich growth regime, Ruddlesden–Popper phases are very unlikely to be formed46. Besides, in resistive switching based on antiphase boundary, the HRS should again show the same resistivity like the initial state, but not a higher one like it is observed for our MOVPE thin films. These unique features of the MOVPE grown SrTiO3 thin films, which are inconsistent with the established models based on oxygen vacancy drift, require further work to be published in a follow up publication.

In-situ scanning transmission electron microscopy (STEM) measurements. To get insight into potential structural modifications that cause resistive switching in our Sr deficient SrTiO3 samples we performed in-situ STEM measurements. For these measurements, a thin lamella with a thickness of ~100 nm is used that is prepared by focused ion beam (FIB). Pt deposition by electron beam evaporation was used for the front side contact. In order to investigate resistive switching behavior of the device under electrical bias, we applied a number of voltage cycles and increased the voltage after each step (±3 V, ±5 V and ±7 V), through which the device was set into the ON or OFF-state, respectively.

Figure 7 shows cross-section STEM-HAADF images of the device after performing the in-situ IV measurements. The Sr deficient MOVPE layer is visible by reduced Z-contrast compared to the stoichiometric substrate, the platinum contact is visible by its grainy structure.

Figure 7a–c show the device after cycling up to 5 V. No indications of filaments or foreign phases are visible in the SrTiO3 films. After applying higher voltages beyond 7 V (Fig. 7d) contrast fluctuations (bright and dark) are visible in the thin SrTiO3 film on the top of the cathode (substrate). Most pronounced is the feature indicated by an arrow, that appears at reduced HAADF intensity and resembles the appearance of conducting filaments as shown by Du et al.47 These in-situ STEM measurements corroborate our finding that the resistive switching in our Sr deficient SrTiO3 MOVPE thin film, which is observed at substantially lower voltages than those required to form filaments, does not involve filament formation. In contrast, our preliminary studies suggest an alternative scenario for resistive switching in Sr deficient films. For instance, Ti-antisite defects have been reported to be mid-gap defect states that induce ferroelectric polarization due to off-center position48,49. The resistive switching could then be based on trap assisted tunneling or ferroelectric switching. Focused investigations to identify the switching mechanism in our devices are ongoing and will be presented elsewhere.

Summary

We have shown that phase-pure stoichiometric and off-stoichiometric SrTiO3 thin films can be grown on (100) SrTiO3:Nb substrates without extended defects by MOVPE. We took advantage of the independent control of the all components in the gas phase together with a high oxygen partial pressure in MOVPE and varied the composition of the gas phase by adjusting the (Sr/Ti)$_{\text{in}}$ precursor ratio. We have achieved Sr deficiency in the growing layer up to 20% without structural degeneration, as validated by TEM and HRXRD. In addition, to an increasing vertical lattice parameter $d_{\perp}$, Sr deficiency increases the permittivity in SrTiO3, and triggers an intrinsic resistive switching effect with an on–off ratio of about 10$^3$ – while no resistive switching was observed in stoichiometric thin films for the same measurement range. IV measurements at 10 K again showed resistive switching for an off-stoichiometric SrTiO3, MOVPE thin film, rendering ionic diffusion processes unlikely, and instead suggesting a defect-based mechanism. We show that the hysteresis evolves without forming pulse or any abrupt threshold. Moreover, the HRS after onset of switching shows a higher resistivity than the layer below the threshold. This suggests that mass transport by ion diffusion does not play an important role in the switching mechanism. Supposingly, in-situ TEM experiments clearly show the absence of filaments in the applied voltage regime.

Methods

SrTiO3 thin films with thicknesses between 40 and 50 nm were epitaxially grown by means of liquid-delivery spin MOVPE50 on 0.5 wt% Nb-doped SrTiO3 (110) substrates (from CrysTec GmbH Berlin, Germany) and DyScO3 (110), both with 0.1° off-cut51. SrTiO3 substrates were pre-treated by etching with a buffered hydrofluoric acid solution and subsequent heating in pure oxygen atmosphere at 1100 °C for one hour. A surface with periodically arranged, atomically smooth ≈ 200 nm wide terraces with single unit cell high steps of ≈ 4 Å (TiO2 termination) are visible in the thin SrTiO3 film on the top of the cathode (substrate). Most pronounced is the feature indicated by an arrow, that appears at reduced HAADF intensity and resembles the appearance of conducting filaments as shown by Du et al.47 These in-situ STEM measurements corroborate our finding that the resistive switching in our Sr deficient SrTiO3 MOVPE thin film, which is observed at substantially lower voltages than those required to form filaments, does not involve filament formation. In contrast, our preliminary studies suggest an alternative scenario for resistive switching in Sr deficient films. For instance, Ti-antisite defects have been reported to be mid-gap defect states that induce ferroelectric polarization due to off-center position48,49. The resistive switching could then be based on trap assisted tunneling or ferroelectric switching. Focused investigations to identify the switching mechanism in our devices are ongoing and will be presented elsewhere.
the total precursor concentration constant. The carrier rotation was adjusted to be 600 rpm to ensure a homogeneous mixture of all gases, a laminar precursor-solvent-O₂/Ar stream and a uniform supersaturation zone above the susceptor. Subsequently, the grown films were cooled down inside the reaction chamber with 10 K/min in 300 mbar O₂ atmosphere by keeping up the carrier rotation constant. The SrTiO₃ films were characterized by HRXRD performed in a Rigaku SmartLab diffractometer using a monochromatic (λ = 1.54056 Å) well collimated primary beam (asymmetric Ge(220) 2-bounce monochromator channel-cut crystal with incidence beam divergence of 12 arc. sec) and a HyPix-3000 horizontal 2D detector. In addition, STEM-HAADF and LAADF was performed in a FEI Titan 80–300 operating at 300 kV, which is equipped with a highly brilliant cathode (X-FEG). The semi-convergence angle was tuned to 9 mrad and the semi-acceptance angle of the detector was set to 35 mrad (camera length 196 mm) and 23 mrad (camera length 300 mm). STEM-EDX was carried out in a JEOL JEM2200FS operated at 200 kV equipped with an X-ray SD detector (Bruker). TEM samples were prepared by plan parallel polishing to a thickness of about 10 µm and final ion milling using a Gatan Pips with liquid nitrogen cooling.

In-situ STEM observations were performed in a same FEI Titan, as for ex-situ measurements above. Cross-sectional TEM specimens from as-grown device was prepared by focused ion beam (FIB) milling using the Thermo Fischer Scientific Nova 600 dual beam system. At around 200 nm thickness, thinning at 30 kV accelerating voltage was stopped to prevent the formation of an amorphous layer on both sidewalls of the specimen. Then, low kV cleaning is carried out at 5 kV and 2 kV to reach the thickness of 100 nm. The FIB lamella was mounted onto the Protochips Fusion 500 double-tilt holder for electrical measurements.

Figure 7. STEM-HAADF images of the device cross-section, after a sweep to (a) −5 V, (b) 0 V, (c) 5 V and (d) 7 V using a current compliance of 30 µA. Dashed lines in (a) indicate the Pt/film/substrate interfaces. Pt layer was deposited on the top of the device to protect the film from ion beam damage during the sample preparation and to serve as an anode for IV measurements.
Electrical and dielectric properties of the homoepitaxial films were investigated at room temperature by preparing the following test structure. Circular platinum contacts of 20 nm thickness with diameters of 150 µm, 200 µm, and 300 µm were deposited through a shadow mask by electron beam evaporation on the as grown films. Thus a MOS test structure composed of a Pt/SrTiOx/SrTiOx:Nb stack was formed. For mechanical contact stability, the platinum was covered with additional 20 nm of nickel. Ohmic backside contacts were formed by liquid Ga/In eutectic. A Keithley 237 source measure unit was used for measuring IV characteristics and by a Boonton capacitance meter 7200 (test frequency 1 MHz, test level 100 mV) the zero-bias capacitance was determined. The measurements were performed in the dark to avoid effects caused by photocurrent or photocapacitance.

Data availability
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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### Author contributions

M.A. and J.S. conceived of the project. A.B., J.S., T.S., K.I., M.A., and J.S. designed experiments. A.B. fabricated the samples and performed X-ray measurements. J.S. measured (di)electric properties. T.S. and T.M. performed electron microscopy. A.B. and C.R. performed X-ray data evaluation. A.B. and W.E. wrote the manuscript. J.M. designed IV experiments at 10 K. H. A. performed in-situ STEM measurements.

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