Modeling precursor diffusion and reaction of atomic layer deposition in porous structures

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I. INTRODUCTION

Atomic layer deposition (ALD) is a vacuum-based technique for depositing thin films of materials with a precise thickness control in the order of less than one nanometer. ALD starts with two gaseous chemicals known as precursors. The solid surface of the substrate is exposed to an alternating sequence of the precursors, separated by purging steps using an inert gas. The first precursor reacts with the surface in a self-limiting manner forming one monolayer (ML). The non-chemisorbed gaseous precursor and the gaseous reaction by-products are removed in a purging and/or an evacuation step. The second precursor also reacts with the surface in a self-limiting manner and thereby re-activates the surface for the first precursor. After a second purging and/or an evacuation step, the surface can be re-exposed to the first precursor. By repeating the cycle of exposition and purging, a thin film is deposited. Due to the self-limitation of the reactions, a maximum of one monolayer of the target material is deposited with each cycle, allowing the thickness of the thin film to be precisely controlled. In consequence of steric hindrance effects or a low density of reactive sites, even less than one monolayer is usually deposited. In this way, inner surfaces of porous structures can also be coated with a low risk of blocking pores.

A wide range of applications exists for ALD, e.g., high-$\kappa$ oxides for CMOS and DRAM technology, solar cell manufacturing, catalysts, and others. In order to improve the control of the deposition process, several models for ALD have been published in recent years. Some of the models use Monte Carlo simulations to describe the ALD process, other models are designed only for high-aspect-ratio structures, partly very complex and partly simplified. Some studies focus only on single parts of ALD, like the sticking coefficient, the growth mode, or the growth per cycle (GPC). A highly useful and practical model, however, was developed by Yanguas-Gil and Elam describing ALD in viscous-flow tubular ALD reactors as well as in porous materials. In their work, transport by viscous flow and diffusion are coupled to surface reactions with first-order irreversible Langmuir behavior. Due to the model’s simplicity, all parameters can be calculated, and the coating result can be predicted for a wide range of precursors and experimental conditions. However, the publication has two disadvantages. The published derivation of the basic differential equations is not comprehensive and elaborated, and an experimental verification of the model for porous materials is missing. In order to overcome these shortcomings, a more detailed, microscopic derivation, based on second-order reaction kinetics, and an experimental comparison with a coated porous substrate is presented in this work. This detailed derivation allows a flexible application of the model to various experimental cases. Thus, it is possible to identify the dependencies of coating results on the microscopic characteristics of the substrate and the precursors used. Additionally, the derivation can be simply adjusted to other features of ALD processes, like different growth modes, precursor depletions, decrease of pore size, and surface diffusion. In order to verify the model, a porous substrate of an anode-supported solid oxide fuel cell (SOFC) was coated and the measured thicknesses of the deposited layer were compared to the predicted coating profile. The aim of the inner.
surface coating of the SOFC substrate is to protect the nickel in the substrate against oxidation. The oxidation of the nickel leads to structural changes of the substrate’s porous microstructure and subsequent to stresses and crack growth in the electrolyte resulting in a complete failure of the SOFC.\textsuperscript{37,38} Thus, the inner surface coating can improve the reoxidation tolerance of anode-supported SOFCs. The study of the influence of ALD coatings on the oxidation tolerance will be presented in another paper.

II. EXPERIMENTAL PROCEDURES

Substrates of anode-supported SOFCs were used as porous structures for the ALD coatings. The substrates were made of a nickel (Ni) and 8 mol% yttria-doped ZrO\textsubscript{2} (8YSZ) cermet and manufactured using the tape casting method (cf. Menzler et al.\textsuperscript{39} for a detailed description).

The ALD ZrO\textsubscript{2} films were deposited in a cylindrical showerhead reactor with a diameter of about 33 cm and a height of about 2.5 cm (LS400C ALD-MOCVD System for Oxides, experimental system for research purposes, supplier VON ARDENNE). ZrO\textsubscript{2} was chosen due to its similarity to cermet and manufactured using the tape casting method (cf. Menzler et al.\textsuperscript{39} for a detailed description).

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The second precursor (O\textsubscript{2}) reacted with all chemisorbed first-precursor molecules (TEMAZ) forming the target material. This assumption was justified because the O\textsubscript{2} pressure in the reactor chamber during O\textsubscript{2} exposure was 80 Pa (only O\textsubscript{2} was present in the reactor chamber), whereas the TEMAZ pressure during TEMAZ exposure was much lower [maximum 1.2 Pa TEMAZ vapor pressure at a bubbler temperature of 30 °C (Ref. 40)]. This means that O\textsubscript{2} diffused deeper into the substrate than TEMAZ and the TEMAZ exposure was the limiting factor for the deposition. The by-products of the chemisorption were assumed to be chemically inert and completely removed from the gas phase during the evacuation/purging step. The initial state of the substrate was assumed to be recovered after one cycle on the deposited layer.

B. Derivation of the differential equations

The transport of the gaseous precursor molecules is described here as pure diffusion, assuming no additional flow inside the pores. Mathematically, the diffusion is described by Fick’s second law\textsuperscript{41}

\[
\frac{\partial n_p(t,z)}{\partial t} = D \frac{\partial^2 n_p(t,z)}{\partial z^2},
\]

where \(n_p(t,z)\) is the volumetric precursor density ([\(n_p\] = m\textsuperscript{-3}) depending on the time \(t\) and the depth \(z\) inside the porous material, and \(D\) is the diffusion coefficient. The precursor molecules in the gas phase react with reactive sites [usually functional hydroxyl groups (OH groups)] on the surface of the pores. Analogous to the TEMAZ/O\textsubscript{2} ALD process, the stoichiometries of each half-reaction of the TEMAZ/O\textsubscript{2} ALD process can be described as\textsuperscript{42}

TEMAZ pulse:

\[
2 \text{surf-OH} + \text{Zr(NR}_1\text{R}_2)_{4(g)} \rightarrow \text{surf-Zr(NR}_1\text{R}_2)_{2(g)} + 2 \text{HN}R_1\text{R}_2_{2(g)},
\]

O\textsubscript{2} pulse:

\[
2 \text{surf-NR}_1\text{R}_2 + x\text{O}_2_{2(g)} \rightarrow 2 \text{Ca}_{a}\text{HbN}_c\text{O}_{d(g)} + 2 \text{surf-OH},
\]
Taking diffusion into account [Eq. (1)], the ALD process is assumed. In this case, the change in the number of surface groups \(N_O(t,z)\) can be written as
\[
\frac{dN_O(t,z)}{dt} = n_O(t,z) \frac{dN_P(t,z)}{dt},
\]
with \(N_p\) as the number of precursor molecules. The precursor density is related to a volume \(V_P\); the density of reactive sites is related to a surface area \(A_O\)
\[
n_P(t,z) = \frac{N_P(t,z)}{V_P},
\]
\[
n_O(t,z) = \frac{N_O(t,z)}{A_O}.
\]
The ratio of \(A_O\) to \(V_P\) is equal to the ratio of the active surface of the porous material and pore volume and is defined as \(\bar{s}\)
\[
\frac{A_O}{V_P} := \bar{s}.
\]
Combining Eqs. (4)–(7), the density change of reactive sites \(n_O(t,z)\) can be written as
\[
\frac{dn_O(t,z)}{dt} = \frac{m}{\bar{s}} \cdot \frac{dN_P(t,z)}{dt} = -\frac{m}{\bar{s}} \cdot n_p(t,z) \cdot n_O(t,z).
\]
Taking diffusion into account [Eq. (1)], the ALD process is then described by two coupled diffusion–reaction differential equations:
\[
\frac{dn_p(t,z)}{dt} = D \frac{d^2n_p(t,z)}{dz^2} - k \cdot n_p(t,z) \cdot n_O(t,z),
\]
\[
\frac{dn_O(t,z)}{dt} = -\frac{m}{\bar{s}} \cdot k \cdot n_p(t,z) \cdot n_O(t,z).
\]
The fraction of available sites \(\Theta(t,z)\) can be defined as
\[
\Theta(t,z) := \frac{n_O(t,z)}{n_O(t=0,z)},
\]
where \(n_O(t=0,z)\) is the density of reactive sites before starting the ALD process, and the degree of surface coverage can be defined as \(1 - \Theta(t,z)\).

C. Determination of the model input parameters

Physically, the change in the fraction of available sites is caused by the number of particles (precursor molecules) sticking on the surface per unit of time and area \(t_p\), normalized to the maximum density of particles, which may adhere to the surface \(\sigma_p\)
\[
\frac{d\Theta(t,z)}{dt} = -\frac{t_p}{\sigma_p} \cdot \Theta(t,z).
\]
The number of particles sticking on the surface per unit of time and area is given by the impinging flux of particles \(J_{wall}\) times the reaction probability \(\beta_0\)
\[
t_p = J_{wall} \cdot \beta_0.
\]
Inserting Eqs. (14) and (13) into (12) and comparing with (10) and (11) yields the reaction rate constant \(k\)
\[
k = \frac{\bar{s}}{m} \cdot \frac{1}{4} v_th n_p(t,z),
\]
where \(v_th\) is the mean thermal velocity, the pressure, and the temperature of the precursor gas above the surface, respectively. \(n_p\) is the mass of a precursor molecule. Inserting Eqs. (14) and (13) into (12) and comparing with (10) and (11) yields the reaction rate constant \(k\)
\[
k = \frac{\bar{s}}{m} \cdot \frac{1}{4} v_th \beta_0,
\]
where \(\beta_0 = 1/\sigma_p\) is the average surface area of an adsorption site. The temperature dependence of the reaction probability \(\beta_0\) is usually assumed to be a Boltzmann factor. However, in this work \(\beta_0\) is constant because the corresponding reaction activation energy and the temperature of the substrate surface do not vary. The maximum density of particles which may adhere to the surface \(\sigma_p\) can be determined from density functional theory by calculating the size of the precursor molecule. Usually, a precursor molecule consists of a central atom (M) and ligands (L). As schematically shown in Fig. 1, large ligands can shield some reactive sites, preventing precursor molecules from reacting, although reactive sites still exist.

The density of (available) reactive sites before the ALD process begins \(n_O(t=0,z)\) is given by
\[
n_O(t=0,z) = m \cdot \sigma_p.
\]
The reason for this is that only reactive sites capable of reacting with the precursor molecules have to be taken into account, but no reactive sites which are shielded by ligands. Inserting Eqs. (16), (15), and (11) into Eqs. (9) and (10) gives the two coupled diffusion–reaction differential equations, which were also reported in Ref. 35:
\[
\frac{dn_p(t,z)}{dt} = D \frac{d^2n_p(t,z)}{dz^2} - \frac{1}{4} v_th \beta_0 \cdot n_p(t,z) \cdot \Theta(t,z),
\]
For low operating pressures (e.g., 80 Pa) during the ALD process, the mean free path of the precursor molecules is much longer than the pore size of the porous material. In this case, the transport of the particles is dominated by Knudsen diffusion. For a porous material with a porosity $\epsilon$, a tortuosity $\tau$, and a mean pore radius $r$, the Knudsen diffusion coefficient is given by

$$D = \frac{2}{\tau^3} \left( \frac{8k_B T_P}{\pi n_P} \right)^{1/2}.$$  

(Take note of the altered exponent for $D$; the 3/2 term is no longer present.)

For high porosities (e.g., 0.61) during the ALD process, the mean free path of the precursor molecules is much shorter than the pore size of the porous material. In this case, the transport of the particles is dominated by Knudsen diffusion. For a porous material with a porosity $\epsilon$, a tortuosity $\tau$, and a mean pore radius $r$, the Knudsen diffusion coefficient is given by

$$\frac{\partial \Theta(t, z)}{\partial t} = -\frac{1}{4}v_{th} \beta_0 \cdot n_p(t, z) \cdot \Theta(t, z).$$  

(18)

For low operating pressures (e.g., 80 Pa) during the ALD process, the mean free path of the precursor molecules is much longer than the pore size of the porous material. In this case, the transport of the particles is dominated by Knudsen diffusion. For a porous material with a porosity $\epsilon$, a tortuosity $\tau$, and a mean pore radius $r$, the Knudsen diffusion coefficient is given by

$$D = \frac{2}{\tau^3} \left( \frac{8k_B T_P}{\pi n_P} \right)^{1/2}.$$  

(19)

Tables I and II list all parameters required to solve the differential equations (17) and (18). The microstructural parameters of the porous material in Table I must be measured for every porous material to be coated.

The parameters in Table II are related to the precursor TEMA. The temperature of the precursor is assumed to be the same as the substrate temperature, since the precursor diffuses inside the substrate. The precursor density outside the porous material $n_P(t, z = 0)$ is given by the vapor pressure of the precursor inside the bubbler [1.2 Pa at 30°C (Ref. 40)]. However, since the degree of saturation of precursor vapor in the carrier gas is usually not known, the vapor pressure can be regarded as an upper limit for the precursor density. The reaction probability $\beta_0$ is a priori unknown and was determined here by fitting the calculated coating profile to experimental data.

### D. Solving the differential equations

The differential Eqs. (17) and (18) were numerically solved using the command “pdsolve” of the computer algebra system “Maple17” [developed by Waterloo Maple Inc. (Maplesoft)]. This command uses finite difference methods with a centered implicit scheme and discretizes time and space. The differential equations were integrated in time with a space step of 2 μm and a time step of 40 ms. For more information see the pdsolve help page of Maple17. The depth $z$ of the porous material was limited to 500 μm, since this is the typical thickness of a tape cast substrate of a SOFC. For $t = 0$, the precursor molecules were located only outside the porous material, assuming an infinite source and therefore a constant density. In the experiments, at $z = 500$ μm, the diffusion of precursor molecules was stopped by a heating plate that served as a substrate support. This meant that the derivative of the precursor density at this point had to be zero. Hence, the initial and boundary conditions can be summarized as

$$\Theta(t = 0, z) = 1, \forall z \in [0, 500 \mu m],$$

(20)

$$n_P(t, z = 0) = 2.9 \times 10^{20} \text{ m}^{-3}, \forall t \geq 0,$$

(21)

$$n_P(t = 0, z) = 0, \forall z \in [0, 500 \mu m],$$

(22)

$$\frac{\partial n_P(t, z)}{\partial z} \bigg|_{z=500 \mu m} = 0, \forall t \geq 0.$$  

(23)

### E. Influence of precursor density and reaction probability on the coating profile

The degree of surface coverage with TEMA molecules after TEMA exposure and before O₂ exposure $1 - \Theta(t, z)$ is shown in Fig. 2 for different precursor densities outside the porous material $n_P(t, z = 0)$ (a) and different reaction probabilities $\beta_0$ (b).

Figure 2(a) shows that the porous substrate is coated deeper for higher precursor densities. Due to the reaction of the gaseous precursor with the surface, the precursor molecules are removed from the gas phase and the density decreases. Other precursor molecules have to diffuse to this point to compensate for the density gradient until the surface is saturated. For higher precursor densities, the surface is saturated quicker, resulting in a deeper coating at the same exposure time.

Figure 2(b) shows that the coating profile is steeper for high reaction probabilities and becomes shallower for lower reaction probabilities. For high reaction probabilities, the reaction of the precursor with the surface is much faster than the
diffusion and all precursor molecules will react with the surface until saturation before continuing to diffuse. For low reaction probabilities, the precursor molecules diffuse past reactive sites and react with the surface at a deeper position inside the porous substrate.

F. Calculation of the layer thickness from the degree of surface coverage

After exposure to the first precursor (here TEMAZ), the surface was exposed to the second precursor (here O2). The oxygen molecules reacted with the ligands of the precursor molecules on the surface. The reacted ligands left into the gas phase and the central atoms with two additional oxygen atoms remained on the surface. It was assumed that all precursor molecules on the surface reacted with O2 and created ZrO2. The previously shielded reactive sites were therefore available again and the degree of surface coverage was reduced by a factor \( \sigma_p/\rho_{\text{reactive}} \), where \( \rho_{\text{reactive}} \) is the density of reactive sites if no sites are shielded.

For convenience, the ZrO2 molecules on the surface were assumed to be cubic. In this case, \( \rho_{\text{reactive}} \) and the thickness of a deposited ML ZrO2 can be calculated from the volume density of the deposited ZrO2 layer \( \rho_{\text{ZrO2}} \):

\[
\rho_{\text{reactive}} = (\rho_{\text{ZrO2}})^{2/3},
\]
\[
d_{\text{ML-ZrO2}} = (\rho_{\text{ZrO2}})^{-1/3}.
\]

The thickness of the deposited ZrO2 layer \( d_{\text{ZrO2}} \) is then given by

\[
d_{\text{ZrO2}}(t, z) = \frac{1 - \Theta(t, z)}{\rho_{\text{reactive}}} \cdot \frac{\sigma_p}{\rho_{\text{ZrO2}}} \cdot \text{degree of coverage after O2 exposure} \times d_{\text{ML-ZrO2}} \cdot \text{number of cycles}
\]
\[
= [1 - \Theta(t, z)] \cdot \frac{\sigma_p}{\rho_{\text{ZrO2}}} \cdot \text{number of cycles}.
\]

The factor \( \sigma_p/\rho_{\text{ZrO2}} \) is equivalent to the GPC of the layer and can be determined experimentally or calculated if the density of the deposited ZrO2 layer \( \rho_{\text{ZrO2}} \) is known.

IV. RESULTS AND DISCUSSION

The thickness of the deposited layer can be calculated from the degree of surface coverage \( 1 - \Theta(t, z) \) using Eq. (26). To compare the predicted coating profile with the experiment, a tape cast substrate of an anode-supported SOFC was coated and the fracture surface was investigated using SEM. A secondary electron SEM micrograph of the fracture surface is shown in Fig. 3 depicting the topography of the sample.

On the top of the micrograph, the outer top surface of the substrate can be seen. Above this surface, the precursor density \( n_p(t, z=0) \) was assumed to be constant and precursor molecules started to diffuse into the porous substrate. This position was defined as depth \( z=0 \); going down into the micrograph, the depth \( z \) increased. The ZrO2 coating can be...
discerned by the different morphologies on the Ni/8YSZ substrate, indicated by the arrows. The red circle shows the point of connection to the next SEM micrograph (Fig. 4).

The ZrO$_2$ coating can also be seen on the Ni/8YSZ substrate in Fig. 4, and the thickness of the ZrO$_2$ coating can be measured with respect to the outer top surface and therefore with respect to the position $z$ inside the porous substrate. The measured ZrO$_2$ thicknesses are shown in Fig. 5 together with two calculated coating profiles.

Different measured ZrO$_2$ thicknesses were found for the same position inside the substrate and vice versa. This scattering can be explained by locally different porousities, tortuosities, pore sizes, and therefore locally different diffusion coefficients as well as by fracturing of the ZrO$_2$ layer, which was not parallel to the fracture surface shown. The coating profiles were calculated using a measured GPC of 0.93 Å/cycle [in agreement with literature values of maximum 1 Å/cycle (Refs. 42 and 46)] and a reaction probability of $\beta_0 = 2 \times 10^{-4}$, fitted to the measured ZrO$_2$ thicknesses. The fitted reaction probability is in the same range as reaction probabilities reported in the literature.\textsuperscript{18,24,47,48} For a precursor density outside the porous material of $n_p(t, z=0) = 2.9 \times 10^{20}$ m$^{-3}$ [TEMAZ vapor pressure of 1.2 Pa at a bubbler temperature of 30°C (Ref. 40)], the calculated coating profile did not fit the measured ZrO$_2$ thicknesses. Instead, the latter can be described by the model using a slightly lower precursor density of $n_p(t, z=0) = 1.0 \times 10^{20}$ m$^{-3}$.

The reasons for this discrepancy could be, on the one hand, an incomplete saturation of precursor vapor in the Ar carrier gas flow of the bubbler, causing a lower precursor density in the reactor chamber, and on the other hand, the transport of the precursor vapor from the showerhead to the porous substrate (distance of about 2.5 cm), inducing a delayed diffusion within the substrate. In order to improve the coating profile prediction, the flow of the precursor vapor in the reactor chamber should be calculated and coupled to the presented diffusion–reaction model. The degree of saturation of precursor vapor in the carrier gas can be determined by weighing the bubbler during the deposition process. In spite of the small discrepancy between model and experiment, the agreement is remarkable considering that only the reaction probability $\beta_0$ was fitted to experimental values.

The model identifies the influence of the precursors used and of the microscopic characteristics of the substrate to be coated on the coating results. For larger precursor ligands (smaller $\sigma_p$), the surface of the substrate is saturated quicker and the GPC is smaller. A high precursor vapor pressure also causes a fast saturation of the surface, whereas a low vapor pressure and a large specific substrate surface makes it difficult to coat the substrate conformally and uniformly in a reasonable period of time. The derivation of the equations can be adjusted to features of any particular ALD process. A term $-k_{\text{depletion}} \cdot n_p(t, z)$ can be added to Eq. (4) to describe a depletion of precursor molecules\textsuperscript{34} (e.g., recombination of atomic precursor species in plasma-supported ALD) and different growth modes can be taken into account in Eq. (11) by local, position-dependent densities of reactive sites before starting the ALD process $n_p(t = 0, z)$. If the deposited coating thickness is significant with respect to the pore sizes, the Knudsen diffusion coefficient will become position-dependent. To take this into account, the term $D \partial^2 n_p(t, z)/\partial z^2$ in Eq. (9) has to be replaced by $\partial/\partial z(D(z)\partial n_p(t, z)/\partial z)$ and the pore size can be reduced in each cycle by the position-dependent layer thickness. A loss of reactive sites on the surface (e.g., due to adsorption of reaction by-products) can be modeled by adding a term $-k_{\text{ads}} \cdot n_{\text{ads}}(t, z)$ to Eq. (10). Surface diffusion of adsorbed precursor molecules can be taken into account by a time-dependent position of reactive sites on the surface, $N_{\text{ads}}(t, z = N_{\text{ads}}(t, z(t))$ and a corresponding surface diffusion coefficient.

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**Fig. 4.** (Color online) Secondary electron SEM micrograph of a fracture surface of a coated substrate. The micrograph is directly connected to Fig. 3 as indicated by the circle.

**Fig. 5.** (Color online) Measured (crosses) and calculated (solid and dashed line) ZrO$_2$ thicknesses for a precursor exposure time of 10 s and 1135 cycles. To calculate the coating profiles, a measured GPC of 0.93 Å/cycle and a fitted reaction probability of $\beta_0 = 2 \times 10^{-4}$ is used. The solid line corresponds to a precursor density of $n_p(t, z=0) = 2.9 \times 10^{20}$ m$^{-3}$ [TEMAZ vapor pressure of 1.2 Pa at 30°C (Ref. 40)], while the dashed line corresponds to a fitted precursor density of $n_p(t, z=0) = 1.0 \times 10^{20}$ m$^{-3}$.\textsuperscript{3}
V. CONCLUSION

A detailed, microscopic derivation of the basic differential equations of the ALD model reported by Yanguas-Gil and Elam34–36 was presented in this work, together with an experimental verification. The thickness of the deposited film can be calculated in relation to the precursor exposure time and the position inside the porous substrate for a wide range of precursors and experimental conditions. The influence of the porous material’s microstructural parameters, e.g., specific surface, porosity, and pore size, and the influence of precursor properties, e.g., ligand size and vapor pressure, on the coating profile were identified, owing to the detailed derivation of the basic equations. Additionally, examples outlined how the derivation can be modified to describe other features of ALD processes. The comparison of the predicted coating profile with the experiment showed good, but improvable, agreement.

Due to the model’s flexible applicability, it is highly practical and can be used to predict ALD coating results for a wide range of applications, such as corrosion resistance, deposition of catalytic material, surface activation by depositing functional groups, and others.

Different precursors and different porous structures must be used for additional verification of the model and to identify its limitations. The model only describes an ideal ALD process without considering desorption of precursor molecules, transport mechanisms other than diffusion inside the pores, or precursor flow in the reactor chamber. Shorter purging and/or evacuation steps will neither change the predicted coating profile nor the deposited layer thickness as long as all nonchemisorbed precursor molecules are removed during these steps. However, an incomplete removal of nonchemisorbed precursor molecules is not described by the model, nor is as an incomplete reaction of O2 with TEMAZ molecules on the surface. In future work, the model should be extended accordingly and verified for systems where these aspects are important.

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1T. Suntola and J. Antson, U.S. patent no. 4,058,430 (15 November 1977).
2T. Suntola and J. Hyvarinen, Annu. Rev. Mater. Sci. 15, 177 (1985).
3S. M. George, A. W. Ott, and J. W. Klaus, J. Phys. Chem. 100, 13121 (1996).
4S. Haukka and T. Suntola, Interface Sci. 5, 119 (1997).
5M. Leskela and M. Ritala, Thin Solid Films 409, 138 (2002).
6M. Ritala and M. Leskela, Handbook of Thin Films Materials: Deposition and Processing of Thin Films, edited by H. Nalwa (Academic, New York, 2002), Vol. 1, pp. 103–159.
7R. L. Puurunen, J. Appl. Phys. 97, 121301 (2005).
8A. Jones, H. Aspinall, P. Chalker, R. Potter, T. Manning, Y. Loo, R. O’Kane, J. Gaskell, and L. Smith, Chem. Vap. Deposition 12, 83 (2006).
9J. Niinistö, K. Kukli, M. Heikkilä, M. Ritala, and M. Leskelä, Adv. Eng. Mater. 11, 223 (2009).
10J. A. van Delft, D. Garcia-Alonso, and W. M. M. Kessels, Semicond. Sci. Technol. 27, 074002 (2012).
11C. Detavernier, J. Dendooven, S. Pulinthanathu Sree, K. F. Ludwig, and J. A. Martens, Chem. Soc. Rev. 40, 5242 (2011).
12F. Zaeera, J. Mater. Chem. 18, 3521 (2008).
13J. R. Bakke, K. L. Pickrath, T. P. Brennan, and S. F. Bent, Nanoscale 3, 3482 (2011).
14J. W. Elam, N. P. Dasgupta, and F. B. Prinz, MRS Bull. 36, 899 (2011).
15Q. Peng, J. S. Lewis, P. G. Hoertz, J. T. Glass, and G. N. Parsons, J. Vac. Sci. Technol., A 30, 010803 (2012).
16H. C. M. Knoop, M. E. Donders, M. C. M. van de Sanden, P. H. L. Notten, and M. W. M. Kessels, J. Vac. Sci. Technol., A 30, 010801 (2012).
17C. Wiemer, L. Lamagna, and M. Fanciulli, Semicond. Sci. Technol. 27, 074013 (2012).
18J. W. Elam, D. Routkevitch, P. P. Mardilovich, and S. M. George, Chem. Mater. 15, 3507 (2003).
19R. A. Adomaitis, J. Cryst. Growth 312, 1449 (2010).
20H. C. M. Knoop, E. Langereis, M. C. M. van de Sanden, and W. M. M. Kessels, J. Electrochem. Soc. 157, G241 (2010).
21M. Rose, Ph.D. thesis (Fakultät Elektrotechnik und Informationstechnik der Technischen Universität Dresden, 2010).
22J. Kim, J. Ahn, S. Kang, and J. Kim, J. Appl. Phys. 101, 073502 (2007).
23A. Lankhorst, B. Pahrazius, H. Terhorst, P. Simons, and C. Kleijn, Surf. Coat. Technol. 201, 8842 (2007).
24R. A. Adomaitis, Chem. Vap. Deposition 17, 353 (2011).
25R. C. Gordon, D. Hausmann, E. Kim, and J. Shepard, Chem. Vap. Deposition 9, 73 (2003).
26J. Dendooven, D. Dedutytsche, J. Mussenhoost, R. L. Vanneirhaeghe, and C. Detavernier, J. Electrochem. Soc. 156, P63 (2009).
27M. Rose and J. Bartha, Appl. Surf. Sci. 255, 6620 (2009).
28M. Rose, J. Bartha, and I. Endler, Appl. Surf. Sci. 256, 3778 (2010).
29R. Puurunen, Chem. Vap. Deposition 9, 249 (2003).
30R. Puurunen, Chem. Vap. Deposition 9, 327 (2003).
31R. Puurunen, Chem. Vap. Deposition 10, 159 (2004).
32R. L. Puurunen et al., J. Appl. Phys. 96, 4878 (2004).
33R. L. Puurunen and W. Vandervorst, J. Appl. Phys. 96, 7686 (2004).
34A. Yanguas-Gil and J. W. Elam, J. Vac. Sci. Technol., A 30, 01A159 (2012).
35A. Yanguas-Gil and J. W. Elam, Chem. Vap. Deposition 18, 46 (2012).
36A. Yanguas-Gil and J. W. Elam, ECS Trans. 41, 169 (2011).
37J. Malzbender, E. Wessel, and R. W. Steinbrech, Solid State Ionics 176, 2201 (2005).
38M. Ettler, H. Timmermann, J. Malzbender, A. Weber, and N. Menzler, J. Power Sources 195, 5452 (2010).
39H. N. Menzler, F. Tietz, S. Uhlenbruck, H. P. Buchkremer, and D. Stöver, J. Mater. Sci. 45, 3109 (2010).
40D. Monnier, I. Nuta, G. Chatillon, M. Gros-Jean, F. Volpi, and E. Blanquet, J. Electrochem. Soc. 156, H71 (2009).
41P. W. Atkins, Physikalische Chemie (Wiley-VCH, Weinheim, 2006).
42W. Weinreich et al., J. Vac. Sci. Technol., A 31, 01A123 (2013).
43T. Zilbauer, Ph.D. thesis (Universität der Bundeswehr München, 2009).
44R. Brodky and H. Hershey, Transport Phenomena: A Unified Approach (McGraw-Hill, New York, 1988).
45W. Schabauer, N. H. Menzler, and H. P. Buchkremer, Int. J. Appl. Ceram. Technol. 11, 125 (2014).
46D. M. Hausmann, E. Kim, J. Becker, and R. G. Gordon, Chem. Mater. 14, 4350 (2002).
47G. Prechtl, A. Kersch, G. Schulze Icking-Konert, W. Jacobs, T. Hecht, H. Boubekeur, and U. Schroder, IEEE Int. Electron Devices Meet., Tech. Dig. 2003, 9.6.1.
48A. Yanguas-Gil and J. W. Elam, J. Vac. Sci. Technol., A 32, 031504 (2014).
49F. Forreiter, Master’s thesis (Karlsruher Institut für Technologie, Institut für Werkstoffe der Elektrotechnik, 2012).
50A. Leonide, Y. Apel, and E. Ivers-Tiffée, ECS Trans. 19, 81 (2009).