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Atomic layer deposition of LiF using $\text{LiN}(\text{SiMe}_3)_2$ and SF_6 plasma†

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Lithium fluoride films were prepared by atomic layer deposition (ALD) using a new route in which $\text{LiN}(\text{SiMe}_3)_2$ is used as a precursor and SF_6 plasma as a coreactant. Conformal LiF films were deposited at 150 °C at a growth rate of ~ 0.4 Å per cycle. All deposited films were polycrystalline and slightly lithium-rich with a composition of $\text{LiF}_{0.8}$, independently of the plasma conditions (e.g. exposure time, pressure and power). The levels of H, C, N, O, Si, and S were all <1 at%. Spectroscopic ellipsometry measurements were carried out over the wavelength range of 140–2480 nm and showed a refractive index of 1.37 at 633 nm for films deposited using 1 s plasma exposure time. We conclude that short plasma exposures are preferred, since a prolonged exposure time leads to an increase in optical absorption and lower growth per cycle values. Furthermore, mass spectrometry measurements revealed the formation of SiMe_3F species during both half-cycles, originating from the reaction between the precursor ligands and fluorine species present either at the surface or in the plasma. Moreover, the SF_6 plasma step led to the formation of fluorocarbon species, suggesting that dissociation and recombination in the plasma takes place. Overall, this work demonstrates that SF_6 plasma offers a promising alternative to other coreactants for ALD of high purity lithium fluoride.

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1. Introduction

Lithium fluoride (LiF) is of large scientific and technological interest because of several emerging applications. LiF films have a large electrochemical stability window of 0–6.4 V vs. Li^+/Li ,⁸ a good chemical stability,^{3,9} and a high mechanical strength.^{4,10,11} LiF has therefore gained attention for its role as an electrode protective/passivating film or as an additive to improve electrode performance in lithium ion batteries.^{4,11,12} Moreover, LiF has one of the largest optical band gaps,¹³ and has a low refractive index of 1.39 at 633 nm.¹⁴ Due to these properties, LiF is useful as a protective transparent optical coating to preserve performance in the ultraviolet region of e.g. aluminum mirrors for space applications.^{7,15} LiF has also been employed as the electron injection layer of organic light-emitting diodes.^{16,17} Moreover, LiF has been explored as a promising electron contact layer in solar cells.^{18,19}

So far, lithium fluoride based thin films have been deposited with various physical vapor deposition methods, including thermal and electron beam evaporation and pulsed laser deposition.^{9,20–22} In the recent years, atomic layer deposition

(ALD) of LiF was demonstrated.^{1–7} ALD is a thin film growth technique based on sequential and self-limiting half reactions. Its benefits of precise thickness control at sub-nanometer level and ability to deposit uniform and conformal films on high-aspect ratio and large-area substrates, are widely recognized.²³ These merits will enable applications of LiF such as 3D-nanostructured batteries or solar cells with a demanding surface topology. Moreover, LiF ALD cycles can be combined with ALD cycles of other materials to tune the film properties. This is the case of LiF combined with AlF_3 to deliver lithium aluminum fluoride, serving as e.g. solid-state Li-ion electrolyte. A literature overview of LiF ALD is presented in Table 1. LiF has for example been deposited using LiTMHD or LiO^tBu as lithium precursor and TiF_4 as coreactant (the latter also in

Table 1 Overview of process details collected from earlier and the present work on the ALD of lithium fluoride

Precursor	Coreactant(s)	T_{sub} (°C)	GPC (Å)	Ref.
LiTMHD ^a	TiF_4	250–350	1.5–1.0	1
LiTMHD ^a	$\text{TiF}_4 + \text{Mg}(\text{TMHD})_2^a$	300–350	1.6–1.4	2
LiO^tBu	TiF_4	200–300	0.35–0.48	3
LiO^tBu	HF-pyridine	150	0.8	4
LiO^tBu	NH_4F	150–300	0.35–0.65	5
$\text{LiN}(\text{SiMe}_3)_2$	HF-pyridine	125–250	0.2–0.6	6
$\text{LiN}(\text{SiMe}_3)_2$	Anhydrous HF	150	0.15	7
$\text{LiN}(\text{SiMe}_3)_2$	SF_6 plasma	150	0.37	This work

^a TMHD = 2,2,6,6-tetramethyl-3,5-heptanedionate.

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combination with $\text{Mg}(\text{TMHD})_2$). Otherwise, anhydrous HF and HF-pyridine have also been adopted as the fluorine source. The combination of LiTMHD with TiF_4 lead to high growth per cycle (GPC) values and nearly stoichiometric films with impurity levels <1 at%.¹ However, the deposition has proven challenging, as the growth was highly dependent on the temperature gradient inside the reactor and required a relatively large LiTMHD dose to deliver uniform films. Furthermore, the films were often characterized by poor adhesion to the substrate. Although these drawbacks can be partially overcome by introducing an additional $\text{Mg}(\text{TMHD})_2$ dose step,² the process becomes complicated. The combination of LiO^tBu and TiF_4 delivered much better film uniformity,³ yet no ALD saturation behavior was reported. More recently, LiF has been deposited using anhydrous HF,⁷ or a HF-pyridine solution,^{4,6} leading to lower deposition temperatures and films with good uniformity. However, the HF precursor is potentially dangerous and corrosive, and although HF-pyridine solutions provide a safer alternative to anhydrous HF and the use of compressed gas cylinders is avoided, strict safety precautions are still required. Very recently, ALD LiF based on a less harmful chemistry, *i.e.* LiO^tBu and NH_4F , has been reported.⁵

In our present work, an SF_6 -fed plasma is exploited as fluorine source for ALD of LiF films. SF_6 plasmas, as well as other F-containing plasmas such as CF_4 and NF_3 , are extensively used for etching of Si-based materials (Si, SiO_2 , Si_3N_4 , *etc.*) and chamber cleaning. For the latter purpose, fluorine plasmas are readily available in many plasma operated ALD reactors. The use of a SF_6 plasma as coreactant has so far only been investigated for the deposition of AlF_3 .²⁴ It was demonstrated that this approach is a promising alternative to the more commonly used anhydrous HF and HF-pyridine as the F-source for ALD of AlF_3 . SF_6 is a stable, non-toxic gas which is easy to handle. Here, we combine SF_6 plasma with $\text{LiN}(\text{SiMe}_3)_2$ as the lithium precursor. $\text{LiN}(\text{SiMe}_3)_2$ has a relatively low melting point of 70°C and it can be used in vapor draw mode. Therefore, the temperature of the precursor delivery line as well as of the walls of the vacuum chamber can be kept relatively low and line clogging is less probable. We examined the growth, film properties as well as surface reactions involved in the LiF plasma-ALD process. ALD behavior was confirmed at 150°C and the growth rate of the films was approximately $0.4 \text{ \AA}$ per cycle. All investigated films were found slightly lithium-rich ($\text{LiF}_{0.8}$) with negligible levels of impurities. From spectroscopic ellipsometry measurements it followed that films were characterized by optical absorption in the shorter wavelength range. The latter increased with the plasma exposure time. To gain more insight into the ALD growth of LiF, quadrupole mass spectrometry (QMS) studies were carried out to identify the reaction mechanisms during the two half-cycles and the similarities and differences with the thermal ALD process of LiF are highlighted.

2. Experimental

The LiF films were fabricated using the Oxford Instruments FlexAL™ thermal and remote plasma ALD reactor. The reactor consists of a rotary and turbo molecular pump such that it can

reach a base pressure of $<10^{-6}$ torr by overnight pumping. The pump unit as well as the inductively coupled plasma (ICP) source are connected to the deposition chamber through gate valves.

In Fig. 1 the ALD process developed in this work is shown. The LiF ALD cycle consists of the exposure of the substrate to $\text{LiN}(\text{SiMe}_3)_2$ (97%, Pegasus Chemicals) as precursor and a SF_6 (99.995%) fed plasma as coreactant. Note that in the gas phase, $\text{LiN}(\text{SiMe}_3)_2$ is mainly present as a dimer but also a small percentage of monomers are observed.²⁵ Because of their lower coordination, monomers are expected to be more reactive than dimers and can dominate the ALD reaction mechanism. In line with previous ALD studies, only monomers are considered in this work.^{6,26} The $\text{LiN}(\text{SiMe}_3)_2$ dose and SF_6 plasma step are alternated by Ar purges to remove all the unreacted species and reaction products from the chamber. The $\text{LiN}(\text{SiMe}_3)_2$ precursor was kept in a stainless steel container and inserted into the chamber by vapor draw for the standard process. Also, the option to bubble the precursor with Ar carrier gas was investigated and is discussed in next section. The $\text{LiN}(\text{SiMe}_3)_2$ precursor pod was kept at 85°C and the supply line was heated to 120°C to prevent precursor condensation. The power for the SF_6 plasma was set to 300 W. The pressure during processing was kept constant at 50 mTorr and a 100 sccm SF_6 gas flow was continuously injected from the top *via* the inductively coupled plasma (ICP) valve. It was confirmed that no film growth occurs without ignition of the plasma, indicating that $\text{LiN}(\text{SiMe}_3)_2$ does not react with SF_6 gas. The wall temperature was set to the maximum temperature of 120°C .

Depositions were performed on a Si(100) wafer with native oxide (Siegert Wafer, 10–20 Ohm cm) and on a Si wafer with micropillars which are $50 \mu\text{m}$ high and have a diameter of $3 \mu\text{m}$ and $5 \mu\text{m}$ spacing. Since SF_6 plasmas can be used for etching of Si-based materials, 10 Al_2O_3 ALD cycles ($\sim 1.5 \text{ nm}$) were used as a protective layer prior to LiF processing. Substrates were placed on a 200 mm carrier wafer and the table temperature was set to 150°C .

In order to obtain the optical properties of the LiF films over the wavelength range of 140–2480 nm, *ex situ* spectroscopic ellipsometry (SE) measurements were performed using vacuum ultraviolet variable angle SE (VUV-VASE) at J.A. Woollam Co., Inc. *Ex situ* measurements were performed at angles of incidence from 55 – 75° with steps of 5° , and photon energy steps of 0.05 eV. The SE model consisted of a combination of one

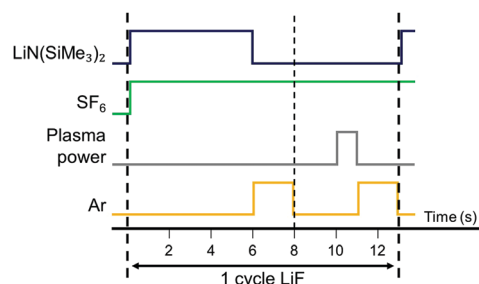


Fig. 1 Schematic of the pulsing sequence of the ALD cycles for LiF.



Sellmeier function and Gaussian oscillator functions. The number of Gaussian oscillators needed to describe the dielectric function increased for longer plasma exposure times. The surface roughness layer was fitted using a Bruggeman Effective Medium Approximation (EMA). Herein, the roughness layer refractive index is described as a mixture of 50% film and 50% voids. *Ex situ* SE was also used to probe thickness uniformity over 200 mm Si wafers. The film thickness was monitored by *in situ* SE with a J.A. Woollam, Inc. M-2000F (250–1000 nm) system. The results from the *ex situ* SE data were used to fit the data obtained *in situ* but without taking the surface roughness layer into account as it lead to correlation between fitting parameters. This simplification hardly affected the variation in thickness which was the only parameter extracted.

Thickness values were confirmed by high resolution SEM (Zeiss Sigma) operated at 2 kV acceleration voltage. An average of at least 5 measurements was taken to determine the uncertainty. SEM was also used to investigate the morphology of the deposited layers. The samples were sputter coated with a 5 nm conductive Au/Pd film prior to the SEM measurements to prevent accumulation of electrons on the sample surface.

The LiF films were also investigated by X-ray photoelectron spectroscopy (XPS).²⁷ XPS spectra were recorded by a Thermo Scientific K-Alpha+ system using monochromatic Al K α X-rays. The spot size of the beam was 400 μ m and the base pressure of the system was 10^{-6} mTorr. Sensitivity values of 0.06 and 4.43 were used for composition analysis of the Li 1s and F 1s regions, respectively. These are the standard values from the database associated with the instrumentation of the Avantage XPS software. The background used to analyze the peaks is a Smart background. The Smart background is an iterative Shirley background to which a constraint has been added such that at no point a background will have a greater intensity than the actual data.²⁸

Elastic backscattering (EBS), elastic recoil detection (ERD) and Rutherford backscattering spectroscopy (RBS) were carried out by Detect99 to determine the composition and mass density of the films. A 2000 keV He⁺ beam was applied for both ERD and RBS. ERD was performed to determine the hydrogen content using 75° sample tilt and the detector at a recoil angle of 25°. For determination of the other elements, RBS was performed with perpendicular incidence and in channeling mode to reduce the background under the Li, C and O features and with two detectors, at scattering angles of 170° and 107°. EBS was necessary for a more accurate determination of the Li content in the films. Here, a 2800 keV proton beam was used in combination with glancing angle geometry to enhance the Li peak area to (Si) background height; also here the sample was tilted by 75°. The scattering angle was 170° in Cornell geometry. For all ion beam techniques, the product of the number of incident particles with the opening angle of the detector has been determined using a beam chopper. The product was calibrated using a reference sample of known composition and thickness. In the case of EBS and RBS, the reference sample was a 100 nm Ni layer on c-Si. For ERD it was a 180 nm Si₃N₄ layer containing 3.4 at% H. For each sample, the two RBS

spectra were simulated simultaneously with the EBS spectrum by NDF (Nuno's DataFurnace), a computer code for ion beam analysis (IBA) data fitting.²⁹ For the EBS simulations, ⁷Li and ¹⁹F cross sections of Paneta *et al.* have been used.³⁰ In the rest of the paper we refer to the combination of RBS, EBS and ERD as IBA. The crystal structure of the layers was determined using a Philips X'Pert MPD diffractometer equipped with a CuK α source (1.54 Å radiation).

A Pfeiffer Vacuum mass spectrometer with a mass-to-charge (*m/z*) range of 100 atomic mass units (amu) was connected to the deposition chamber through a pipeline and a 150 μ m diameter pinhole. The system is equipped with a Channeltron detector and the energy of the electrons in the ionizer was set to 70 eV. The pressure in the QMS was maintained below 10^{-6} torr using differential pumping with a turbomolecular pump. Prior to every measurement the background signal and pressure of the reactor were measured as a reference. The measuring time per atomic mass unit (amu) was set to 200 ms. For the time-resolved measurements, the selected *m/z* values were tracked per channel using a dwell time of 50 ms. The QMS data of at least 10 cycles were monitored to verify that steady-state values were achieved. The measurement procedure was explained more extensively in our previous work.³¹

3. Results and discussion

3.1 ALD process

Film growth of ALD LiF was examined by *in situ* SE using a deposition temperature of 150 °C. In Fig. 2 the film thickness is shown as a function of the number of cycles. The first 50 cycles show a slightly enhanced growth compared to the steady growth regime. This can be due to an initial growth effect, but it can also originate from the fitting procedure to extract the thickness from the ellipsometry data. In the fit, the optical properties of the film have been assumed constant throughout the whole thickness but this assumption is not necessarily accurate. The growth per cycle is ~ 0.4 Å, which is in the range of reported literature values for processes which use LiN(SiMe₃)₂ as the lithium precursor.^{6,7} The growth was also monitored after each half-cycle, as shown in the inset of Fig. 2. The magnitude of the changes in this “apparent” thickness after the precursor and SF₆ plasma steps do not necessarily reflect the real change in physical thickness because the top surface is not included in the model. The inset shows that, after the LiN(SiMe₃)₂ dosing, the apparent thickness increases by several angstroms, because of the chemisorption of the precursor on the surface. After the SF₆ plasma step, the apparent thickness decreases because of the removal of the organic ligands. It can be expected that the precursor ligands are abstracted and the surface is then terminated with fluorine. The reaction mechanism is further discussed in Section 3.3.

In Fig. 3, the saturation curves for the ALD process of LiF are shown. The growth per cycle was determined as a function of the LiN(SiMe₃)₂ dosing time, SF₆ plasma exposure time and purge times. The saturation of the pressure during the



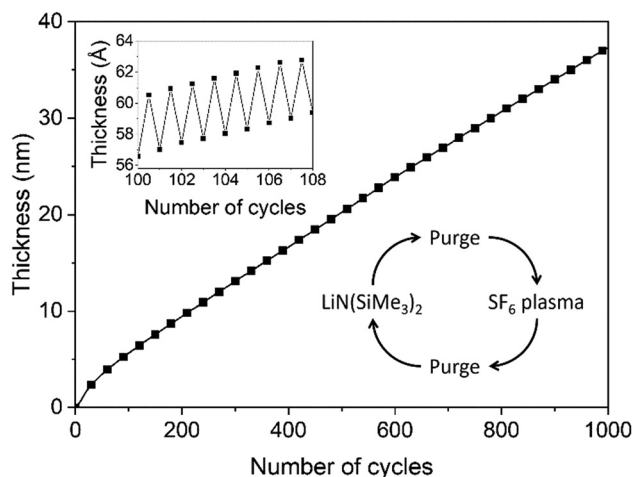


Fig. 2 Film thickness as a function of the number of cycles measured by *in situ* spectroscopic ellipsometry for ALD of lithium fluoride using a 6 s $\text{LiN}(\text{SiMe}_3)_2$ dose and 1 s plasma exposure at 150 °C. The inset shows the apparent thickness measured every half-cycle.

$\text{LiN}(\text{SiMe}_3)_2$ dose and plasma power are reported in Fig. S1 (ESI†). To extract the growth per cycle, the film thickness was monitored every 10 cycles while changing the dose time of the reagents, pressure or argon purge. The self-limiting behavior of the $\text{LiN}(\text{SiMe}_3)_2$ dosing time was tested with and without the use of an Ar carrier gas (Fig. 3a). In some cases, the use of an Ar carrier gas could lead to the decrease of the precursor saturation time. However, no difference was observed for our process, meaning that the vapor pressure of the precursor is high enough to dose a sufficient amount of precursor in the reactor and avoid the use of a carrier gas. The growth per cycle *versus* $\text{LiN}(\text{SiMe}_3)_2$ dosing time shows a so-called soft saturation behavior. A $\text{LiN}(\text{SiMe}_3)_2$ dosing time of 6 s was selected for the recipe. A purge step of 2 s was enough to remove the reaction products and unreacted precursor molecules from the chamber. Extending the purge time did not change the growth per cycle value. Fig. 3b reports the growth per cycle as a function of the plasma exposure time. After an initial increase, the growth per cycle decreases again for extended plasma exposure times. Such a particular trend, which was confirmed by SEM analysis (the corresponding SE measurements and SEM images can be found in Fig. S2 and S3, respectively, ESI†) has been observed more often for plasma-ALD processes.³² It can be attributed to several potential causes such as an etching effect playing a role for long plasma exposure times or inhibition of precursor adsorption after exposing the surface for a longer time to F radicals. Another likely cause is that the film properties are affected by the length of the plasma exposure time which has subsequently an effect on the growth itself. As will be addressed in the next section, the optical absorption of the films changes quite significantly for long exposure times showing that the film properties are affected. For the standard recipe, 1 s plasma exposure was selected. Also for the plasma half-cycle a purge step of 2 s was sufficient to remove the reaction products and unreacted molecules from the chamber. Relatively short cycle times of 13 s were defined for the standard process of LiF ALD

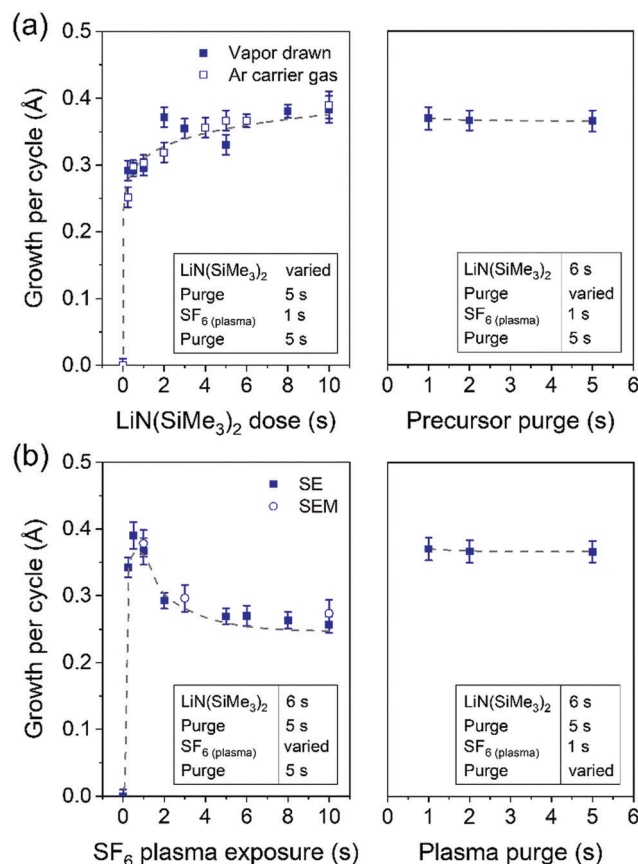


Fig. 3 Growth per cycle obtained by SE as a function of (a) precursor dose and purge and (b) plasma exposure and purge for ALD of LiF obtained at 150 °C. The precursor dose was investigated with and without the use of Ar carrier gas. The SE data in (b) were compared to SEM data (diamond symbols). The dotted lines serve as a guide to the eye.

films (Fig. 1). It is worth reporting that the effect of pressure during the precursor and plasma exposure steps and the effect of plasma power were also investigated. A pressure increase above 50 mTorr during precursor exposure did not lead to any increase in growth per cycle (Fig. S1, ESI†). Moreover, the SF_6 plasma was only stable when the pressure was 50 mTorr or higher. Therefore, a pressure of 50 mTorr was selected for the process. Adjusting the plasma power did not affect the growth per cycle (Fig. S1, ESI†).

To further demonstrate the ALD behavior of the LiF process, the uniformity and conformality were investigated. For both studies the standard recipe was used. Film thickness and refractive index uniformity were determined for a ~90 nm thick film using *ex situ* SE on various points of films deposited on a 200 mm wafer. The measured thicknesses and refractive indices are presented in Table S1 (ESI†). The non-uniformity (standard deviation, 1 sigma) of the thickness and refractive indices were determined to be 4.4% and 0.2%, respectively. This points to rather good uniformity of film thickness and excellent uniformity of the refractive index. However, the process parameters were optimized for the growth per cycle value and not for film uniformity. Therefore, the thickness



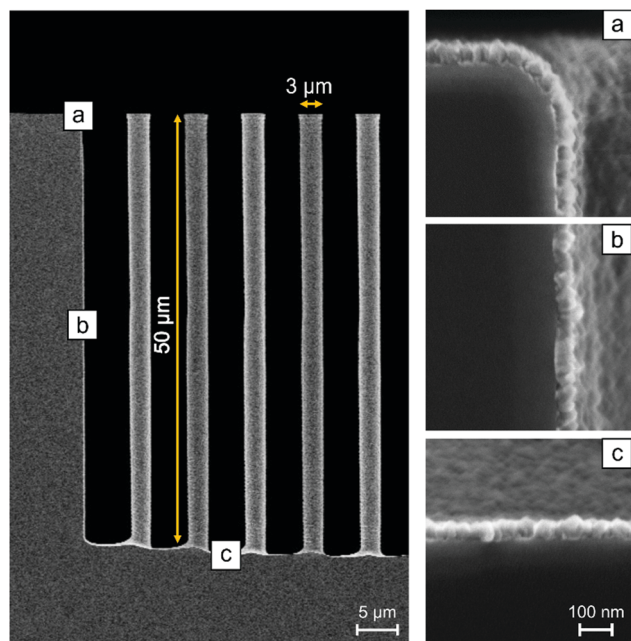


Fig. 4 Analysis of the conformality of the LiF ALD process. Micropillars of 50 μm high and a diameter of 3 μm are used for this purpose. On top of the pillars a 60 nm layer of LiF was deposited. Images (a), (b) and (c) show the LiF layer at respectively the top, middle and bottom of the pillars. Thickness variations from 57–62 nm were observed. The scale bar of 100 nm applies to all insets.

uniformity can probably still be improved. The conformality of the deposition was studied using SEM at different positions on high aspect ratio Si pillars. The Si pillars were covered with a LiF layer of approximately 60 nm. Fig. 4 shows 3 SEM pictures taken at different heights along the pillar. Image (a) is taken at the top of the pillars and shows a LiF layer of 62 ± 2 nm. Image (b) is taken at half-length of the pillar and depicts a 60 ± 2 nm LiF layer (96% of thickness at top). At the bottom (c), the LiF film is 57 ± 2 nm thick (93%). From these measurements we can conclude that a conformal coating on high aspect ratio pillars was achieved.

3.2 Film properties

3.2.1 Chemical composition and mass density. XPS measurements were carried out to analyze the chemical

composition of the LiF films. A survey spectrum can be found in the ESI† (Fig. S4). In Fig. 5, the XPS core level spectra of C 1s, Li 1s, and F 1s are shown for 40 nm thick films prepared at 150 $^{\circ}\text{C}$ using 1 s plasma exposure time. The spectra for different plasma exposures are very similar and therefore not shown here. The binding energy scale was calibrated using the C 1s peak at 284.8 eV, associated with adventitious carbon on the sample surface. The Li 1s peak and F 1s peak are observed at 55.5 eV and 685.1 eV, respectively, both in close agreement with literature reports for LiF films.³³ A sputtering procedure was adopted to investigate the stoichiometry of the film. In contrast to our previous XPS studies of Li_2CO_3 ,³⁴ the stoichiometry of the films did not change during depth profiling. The obtained LiF stoichiometry of the film is 54.4 at% Li, 44.3 at% F, 0.3 at% C and 0.4 at% O. Hence, the films deposited in this work are found to be lithium-rich with a stoichiometry of $\text{LiF}_{0.8}$. The obtained stoichiometry falls in the range of values reported for ALD LiF films, which have a F/Li ratio of 0.73 to 1.23.^{1,2,4–7} Carbon and oxygen were only found at the surface of the films. The atomic percentages of Si, N and S were below the detection limit of the XPS. Literature has also reported the absence of impurities originating from the selected precursor or coreactant. However, in some cases the presence of traces of oxygen or hydrogen was reported.^{3,4} In addition, LiF films deposited using longer plasma exposure times were probed by XPS, since these films differ in terms of film growth as shown in the previous section. It was found that the stoichiometry of the films is rather independent of plasma exposure time. These results and a comparison of the stoichiometry of the films for surface scans and during depth profiling can be found in Table S2 in the ESI†. The influence of pressure and plasma power on the film stoichiometry were also investigated, but no changes in stoichiometry were observed (not shown here). Moreover, film composition did not show noticeable changes after storage in atmosphere for one month (see ESI† Fig. S5). This also holds for the film thickness, as well as the mass density and roughness which will be discussed later. These results suggest that LiF grown by ALD is stable under ambient conditions.

A combination of ion beam techniques, which are nondestructive and give accurate quantitative information, was used to complement the aforementioned XPS data.³⁵ In addition, these

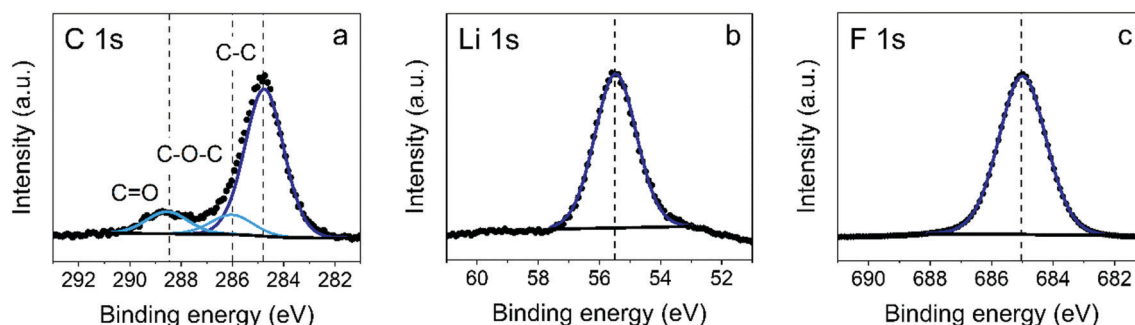


Fig. 5 (a) C 1s, (b) Li 1s, and (c) F 1s spectra of a 40 nm thick lithium fluoride film grown at 150 $^{\circ}\text{C}$ using 1 s plasma exposure time. The measured data (black dotted lines) and fitted peaks (blue lines) are reported. Carbon is detected only at the surface of the films.



techniques allow to probe also the hydrogen content in the film. LiF films deposited using different SF_6 plasma exposure times were studied and the results are shown in Table 2. The original experimental and simulated data are reported in the ESI† (Fig. S6). The films are lithium rich and of high purity, in agreement with previous XPS studies. The IBA data also show that the total number of deposited atoms in a nm^2 per cycle decreases for longer plasma exposure times. Previously, in Fig. 3b, it was shown that the growth per cycle in terms of thickness also decreases for longer plasma exposure times. This means that the thickness decrease observed for longer plasma exposure times is caused by an actual decrease in deposited LiF per cycle. According to the trend in the SE and SEM data, we would expect a further decrease in growth per cycle when prolonging the plasma exposure to 3 and 10 s. From the IBA data this decrease is not confirmed.

A combination of the IBA and SE data was used to calculate the mass density of the samples. From IBA measurements the areal number density of a specific atom (atoms per nm^2) was extracted. The areal densities were multiplied with the mass of the atoms under consideration, and after summation the mass density was obtained by dividing by the film thickness. The mass density for all LiF films was found to be $\sim 2.3 \text{ g cm}^{-3}$. The bulk value of the mass density of stoichiometric LiF reported in the literature is 2.6 g cm^{-3} .³⁶ Typically, thin films prepared by ALD or other techniques are reported to have a somewhat lower mass density compared to the bulk material. Differences in mass density probably also exist due to the sub-stoichiometric nature measured for our LiF films.

3.2.2 Crystallinity and morphology. The deposited films were studied by XRD for various plasma exposure times. Fig. 6 shows the X-ray diffractograms of $\sim 40 \text{ nm}$ thick films obtained using a θ - 2θ Bragg-Brentano geometry. The θ - 2θ measurements, which detect periodicity of planes parallel to the surface, show only the typical diffraction peak of crystalline LiF at 38.7° .³⁷ The small peak near 44° originates from the sample holder. This means that the process yields a preferential crystal growth orientation in the $\langle 111 \rangle$ direction. The FWHM and intensity of the peak at 38.7° do not change for different plasma exposure times within the measurement error. This means that film crystallinity does not significantly depend on the plasma exposure time. The LiF films prepared by thermal ALD reported in the literature also showed crystalline features, although in most studies no preferential growth orientation was reported.^{1–4,6,7} There is only one study on thermal ALD LiF reporting on a preferential crystal growth orientation in the $\langle 200 \rangle$ direction.⁵ The origin of the preferential crystal growth orientation and therefore also the difference between the

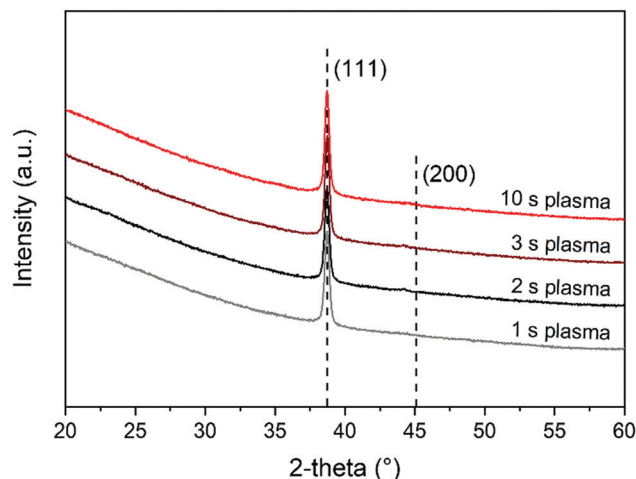


Fig. 6 XRD θ - 2θ scans of $\sim 40 \text{ nm}$ thick LiF films grown on Si. LiF films are deposited using plasma exposure times of 1, 2, 3 and 10 seconds. The dotted lines indicate the positions of the expected diffraction for LiF powder samples.

preferential crystal growth orientation of the thermal LiF process and our plasma based process is unclear.

Fig. 7 shows high resolution SEM images of a 70 nm LiF film deposited using a 1 s plasma dose on top of a Si substrate. The crystallites exhibit a size comparable to the sample thickness and seem to have a triangular shape. Longer plasma exposure times lead to similar results. From the height difference observed in the cross sectional image in Fig. 7b, it can be concluded that there is significant surface roughness. The presence of surface roughness was confirmed by *ex situ* SE measurements. The surface roughness was relatively high, being $\sim 10 \text{ nm}$ for each of the films.

3.2.3 Optical properties. Fig. 8 shows the optical properties as obtained from vacuum ultraviolet (VUV) SE measurements for LiF films deposited at 150°C using different plasma dosing times. As shown in Fig. 8a, the refractive index (n) is 1.37 at 633 nm and shows a relatively low dispersion for the film deposited using 1 s plasma exposure. At 633 nm the refractive indices are 1.38 and 1.39 for 3 s and 10 s plasma exposure, respectively. The refractive index values for LiF films prepared by thermal ALD using HF and TiF_4 as coreactants vary from 1.37–1.39 and are thus in agreement with our findings.^{1,4,6}

As observed from Fig. 8b, absorption occurs in a rather broad range of wavelength. However, LiF films are well known for their large optical band gap and high transparency. The LiF films deposited in this work are slightly sub-stoichiometric ($\text{LiF}_{0.8}$). Anionic vacancies can lead to the formation of lattice defects.

Table 2 Properties of 40 nm thick LiF films prepared using plasma exposure times of 1, 3 and 10 s. The stoichiometry was calculated from IBA results. The mass density was obtained by combining IBA and SE results. In the first row the typical error is given for a certain parameter

Sample	Li at per nm^2 per cycle	F at per nm^2 per cycle	F/Li	[H] (at%)	[O] (at%)	[C] (at%)	Density (g cm^{-3})
1 s plasma	2.0 ± 0.1	1.6 ± 0.1	0.78	1.0 ± 0.1	0.8 ± 0.1	0.1 ± 0.1	2.3 ± 0.1
3 s plasma	1.6	1.3	0.83	0.8	1.1	0.9	2.3
10 s plasma	1.6	1.2	0.75	1.1	0.9	1.3	2.3



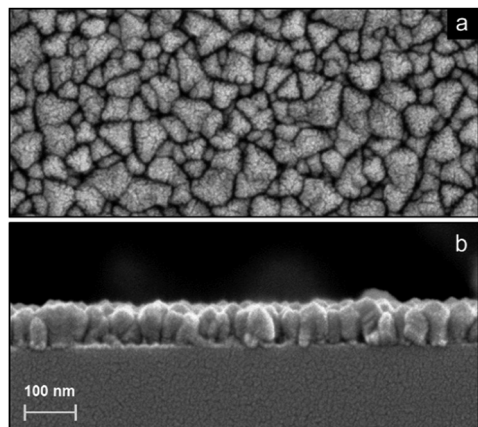


Fig. 7 (a) High resolution SEM images showing the surface topography and (b) cross-section of a LiF film deposited on Si using a 1 s SF_6 plasma exposure. The scale bar applies to both images.

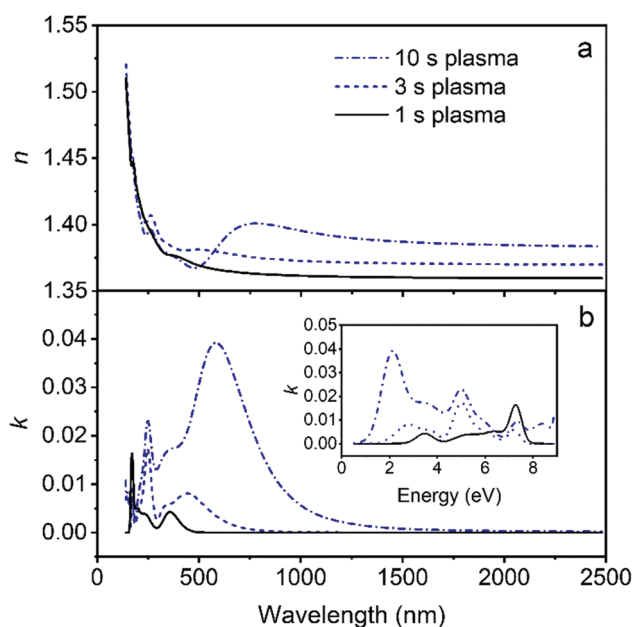


Fig. 8 Refractive index (n) and extinction coefficient (k) as a function of wavelength as determined from VUV-SE for 40 nm thick films deposited using 1, 3 and 10 s SF_6 plasma exposure. The inset shows the extinction coefficient as a function of photon energy.

DFT studies in the literature show that when F vacancies occur, new energy levels develop which may affect the band-gap.³⁸ However, valence band XPS spectrum analysis (Fig. S7, ESI†) shows features at 29–30 eV and 8–10 eV, independent of the plasma exposure time, and in agreement with literature of stoichiometric LiF.³⁹ It should be noted that the band gap itself cannot be determined from the VUV-SE data since it lies outside the measured energy range of 8.8 eV. Finally, we consider the possibility that the plasma environment can play a role in the formation of this absorption features. Specifically, it has been shown that when LiF is irradiated with ions,^{40,41} electrons,⁴² or even (V)UV

radiation emitted from discharges,⁴³ color centers can be produced. The most dominant defects which can be created in LiF are F centers (an electron localized at a fluorine vacancy).⁴¹ As observed from the absorption spectra Fig. 8b, the absorption features become dominant and extend to longer wavelengths upon increasing the SF_6 plasma exposure time from 1 to 10 s. Since the film stoichiometry is the same, independently of the plasma exposure time, we consider it plausible to attribute the presence of the absorption features to changes caused by plasma radiation. This conclusion is also supported by the fact that no absorption features are reported for thermal ALD of LiF.^{1–3,6} Instead, literature studies of AlF_3 films prepared by ALD using an SF_6 plasma also show absorption features in the lower wavelength range.²⁴

3.3 Surface reactions

The gas phase reaction products were analyzed by mass spectrometry to gain insight into the surface reactions occurring during the ALD process. Specifically, we would like to highlight the similarities and differences between the thermal ALD process when using HF, and our SF_6 plasma based LiF ALD process. A study on the surface reactions of thermal ALD of LiF with $\text{LiN}(\text{SiMe}_3)_2$ as precursor and HF-pyridine as coreactant was already presented in the literature.⁶ Herein, it was reported that $\text{LiN}(\text{SiMe}_3)_2$ molecules can either physisorb or chemisorb during the first half-cycle. Physisorption of $\text{LiN}(\text{SiMe}_3)_2$ on LiF was attributed to strong interactions of Si and N in $\text{LiN}(\text{SiMe}_3)_2$ with F and Li in LiF, respectively. During the second half-cycle, the surface is exposed to HF to promote ligand abstraction from the physisorbed $\text{Li-N}(\text{SiMe}_3)_2$ molecules and form LiF. Specifically, the precursor ligands on the surface were abstracted in the form of SiMe_3F and NH_3 . However, it was reported that if physisorbed HF is present, the precursor ligand could instead chemisorb and already lose its ligands, *i.e.* LiF would be formed already during the precursor dosing step.^{6,44} Compared to other metal fluorides, which are shown to adsorb a significant amount of HF after HF exposure, the amount of adsorbed HF on LiF is expected to be limited because of the Lewis base nature of LiF.⁴⁴ LiF would rather act as a F^- ion donor. Despite the presence of peaks at $m/z = 77$ (SiMe_2F^+ from SiMe_3F) and $m/z = 17$ (NH_3^+ from NH_3) in the mass spectrometer at the end of the precursor step, the authors concluded with a complementary quartz crystal microbalance (QCM) study that the precursor ligands are indeed primarily abstracted during the HF exposure. Moreover, they concluded that at higher deposition temperatures ($>200^\circ\text{C}$) the ligand abstraction during the precursor step is suppressed because of the limited physisorbed HF on the LiF surface.

To gain insight into the species present in the reactor during our plasma-ALD process, the spectra during precursor dosing and SF_6 plasma ignition were measured in the range of $m/z = 1$ –100 as shown in Fig. 9. In Fig. 9a the cracking pattern of the $\text{LiN}(\text{SiMe}_3)_2$ precursor is shown. Note that the peak from the parent molecule itself and the peaks with the highest expected intensity (146, 130, and 147 amu, respectively) could not be measured since they lie outside the range of the mass spectrometer.^{45,46} The peak with the highest intensity within



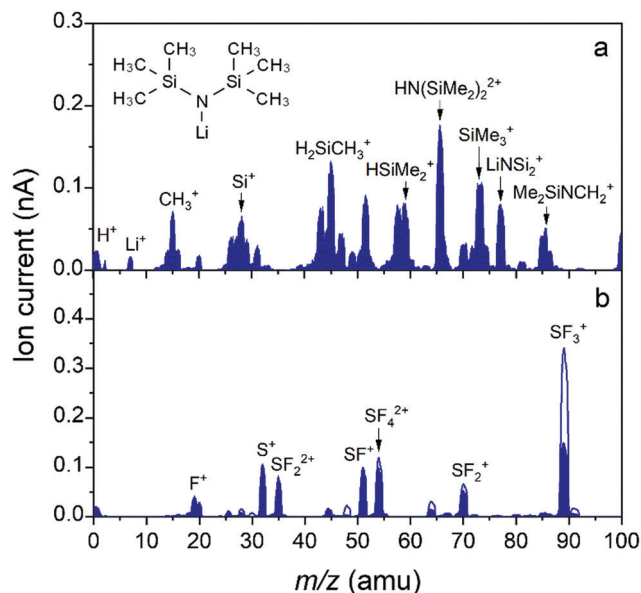


Fig. 9 Ionization pattern obtained by mass spectrometry during (a) $\text{LiN}(\text{SiMe}_3)_2$ precursor dosing, and (b) SF_6 plasma ignition (filled curves). In (a) also the $\text{LiN}(\text{SiMe}_3)_2$ precursor molecule is depicted. The ionization pattern for the SF_6 gas (plasma off) is also added (open curve).

the explored mass range is located at 65.5 amu, and its width would suggest there are multiple contributions to it.^{26,45} We expect the m/z of 65 amu corresponds with the double ionized species from $m/z = 130$ ($\text{N}(\text{SiMe}_2)_2^+$), and the m/z of 65.5 corresponds with the double ionized species from $m/z = 131$ ($\text{HN}(\text{SiMe}_2)_2^+$).^{26,47} Fig. 9b shows the spectra of SF_6 plasma and SF_6 gas, which are essentially identical in terms of detected species. As expected, the peaks during SF_6 plasma exposure are lower than the SF_6 gas spectrum due to dissociation in the plasma. All measured species are ion fragments of SF_6 . Note that again the parent molecule itself (SF_6) could not be measured since it lies at $m/z = 146$. Instead, SF_3^+ at $m/z = 89$ was tracked to measure SF_6 .

A complete overview of the measured species is presented in Table S3 in the ESI† Fig. 10 presents a selection of these species, which are followed during one cycle of LiF. In Fig. 10a–e, the following masses are considered: $m/z = 17$ (NH_3^+ from NH_3), 65.5 (from $\text{LiN}(\text{SiMe}_3)_2$ or $\text{HN}(\text{SiMe}_3)_2$), 69 (CF_3^+ from CF_4 or CHF_3), 77 (SiMe_2F^+ from SiMe_3F or SiMe_2F_2 and LiNSi_2^+ from $\text{LiN}(\text{SiMe}_3)_2$) and 85 (SiF_3^+ from SiF_4 or SiMe_3F and $\text{Me}_2\text{SiNCH}^+$ from $\text{LiN}(\text{SiMe}_3)_2$). Note that $m/z = 77$ is the largest peak in the mass spectrum of SiMe_3F , whereas the peak at $m/z = 92$ (SiMe_3F^+) has negligible intensity.⁴⁶ Additional mass-to-charge ratios 15 (CH_3^+), 31 (CF^+), 33 (CH_2F^+ or SH^+), 34 (CH_3F^+ or H_2S^+), 50 (CF_2^+), 51 (CHF_2^+), and 81 (SiMe_2F^+ from SiMe_3F or SiMe_2F_2) amu can be found in the ESI† in Fig. S8. Also, the pressure in the system was monitored and reported in Fig. 10f. The graph is divided into several sections representing the different ALD process steps, i.e., $\text{LiN}(\text{SiMe}_3)_2$ dose and SF_6 plasma exposure, which are alternated by Ar purging. Compared to the ALD recipe for LiF described earlier, all purge steps are now extended to 5 s to clearly separate the process steps and

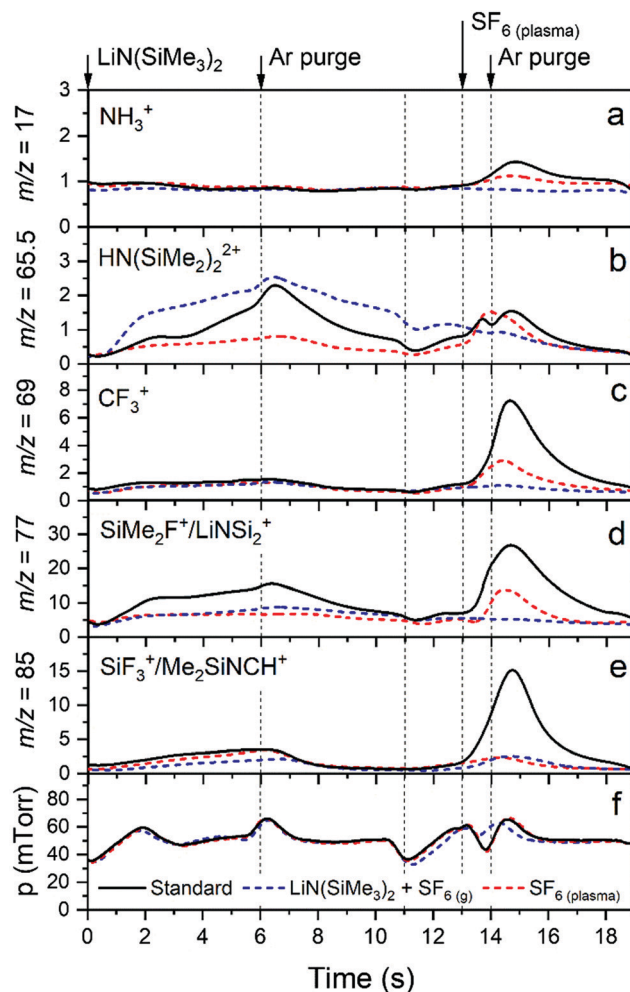


Fig. 10 Time-resolved QMS data of $m/z = 17$, 65.5, 69, 77 and 85 amu after 10 cycles. The standard ALD recipe (black) is compared to the $\text{LiN}(\text{SiMe}_3)_2 + \text{SF}_6$ recipe (blue dotted) and SF_6 plasma recipe (red dotted). The precursor and coreactant dosing steps are alternated by argon purge steps of 5 s.

facilitate data interpretation. (As shown in Fig. 3, extended purge times did not influence the process.) In addition to the standard ALD recipe, in Fig. 10 the data are also reported for recipes where one of the reagents is omitted from the recipe. For these recipes no ALD growth occurred and the data is reported after 10 cycles so that steady-state values were achieved. Therefore, it is possible to distinguish the reaction products from other species present in the reactor and fragmentation and ionization occurring in the mass spectrometer. In every plot three different curves are shown, corresponding with the following recipes: the standard recipe, the recipe with only $\text{LiN}(\text{SiMe}_3)_2$ and SF_6 gas (no plasma), and the recipe with only SF_6 plasma.

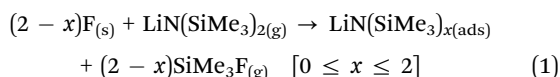
In the first half-cycle the lithium precursor was introduced in the chamber which led to an increase in signal at $m/z = 65.5$ for the standard recipe as well as for the $\text{LiN}(\text{SiMe}_3)_2 + \text{SF}_6(\text{g})$ recipe, as expected. The peak at $m/z = 65.5$ is smaller in the case of the ALD process compared to the $\text{LiN}(\text{SiMe}_3)_2 + \text{SF}_6(\text{g})$ recipe. We attribute this observation to precursor adsorption and



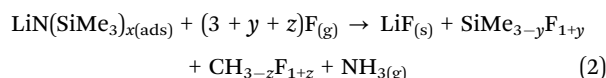
reaction with the surface in the case of the ALD process, whereas for the $\text{LiN}(\text{SiMe}_3)_2 + \text{SF}_6(\text{g})$ recipe the surface was already saturated with Li-precursor. Also a signal at $m/z = 77$ was observed during the $\text{LiN}(\text{SiMe}_3)_2$ dose. The standard ALD recipe has a higher signal compared to the other recipes, pointing out that this is a reaction product of the Li-precursor with the surface, promoted by the presence of F^- species on the LiF surface. As mentioned before, this reaction product was also found in the mass spectrometer for the thermal ALD LiF process at 150°C , although in combination with NH_3 .⁶

In the second half-cycle the surface is exposed to SF_6 plasma. Note that from the QMS measurements no clear evidence of sulfur-containing reaction products was observed. The signal at $m/z = 34$ (Fig. S8, ESI†) could however also originate from H_2S species formed in the plasma. Furthermore, no S is detected in the film. The dominant neutral species in an inductively coupled SF_6 plasma are known to be SF_6 , F, F_2 , and SF_4 , while the dominant ions are SF_5^+ and F^+ .^{48–50} The concentration of S and S^+ is typically a factor 10^3 lower than the concentration of F and F^+ .⁵⁰ Moreover, it is known that F is very reactive due to its very high electronegativity and electron affinity.⁵¹ Therefore, it is reasonable to assume that only F takes part in the surface reactions. During this half-cycle again peaks at $m/z = 65.5$ and 77 were observed, as well as a peak at $m/z = 17$ (NH_3^+). Also other fluoromethylsilane species and fluorocarbons were found. Specifically, reaction products at $m/z = 85$ (SiF_3^+ from SiF_4 or SiMeF_3) and $m/z = 69$ (CF_3^+ from CF_4 or CHF_3) were observed. As shown in the Fig. S8 (ESI†), reaction products at $m/z = 81$ (SiMeF_2^+ from SiMeF_3 or SiMe_2F_2), $m/z = 50$ (CF_2^+) and $m/z = 31$ (CF^+) were observed. Moreover, relatively small peaks at $m/z = 33$ and 34 assigned to CH_2F^+ and CH_3F^+ were found. It can be observed that all masses reported in Fig. 10 (except 65.5 amu, which corresponds with the precursor molecule) show a higher intensity during the second half-cycle with respect to the first half-cycle. This indicates that ALD surface reaction products are primarily generated during the plasma exposure.

To summarize the findings from the QMS studies in the first and second half-cycles: a fraction of the precursor molecules that adsorb at the surface already loses one of its ligands during the $\text{LiN}(\text{SiMe}_3)_2$ dose step. The remaining Li-precursor ligands are abstracted during the plasma step. Moreover, the reaction products can be further dissociated in the plasma leading to *e.g.* fluorocarbon species. Reaction eqn (1) is proposed for the precursor dose, where the adsorbed precursor can undergo partial loss of ligands.



During the SF_6 plasma exposure step, the fluorine plasma species remove the remaining precursor ligands to form LiF. The equation is unbalanced for simplicity as the surface species and the ratio of the reaction products are unknown at this point:



Our results seem to differ from those of thermal ALD of LiF reported in literature. Unlike thermal ALD, a reasonable fraction of the precursor molecules already loses ligands during the first half-cycle. Moreover, in the case of thermal ALD, the reaction during the first half-cycle was explained by the presence of surface adsorbed HF. As mentioned before, LiF is a Lewis base that can share the lone pairs on F. Therefore, we consider it more likely that F from LiF would be partly consumed in the ALD reaction during the first half-cycle. This could also explain why our films are slightly F-deficient. Moreover, unlike thermal ALD LiF, additional fluoromethylsilane and fluorocarbons are detected in the second half-cycle. Most likely dissociation and recombination in the SF_6 plasma takes place.

4. Conclusions

A new ALD chemistry has been demonstrated to prepare LiF films using $\text{LiN}(\text{SiMe}_3)_2$ and SF_6 plasma. Self-limiting growth was obtained at ~ 0.4 Å per cycle and conformal coverage of high aspect ratio pillars was achieved for deposition at 150°C . Moreover, high purity films were obtained with contamination levels < 1 at%. The obtained stoichiometry of the films was $\text{LiF}_{0.8}$ as determined by IBA and XPS measurements. Furthermore, the ALD process yields polycrystalline films with preferential growth in the $\langle 111 \rangle$ direction. The film density was 2.3 g cm^{-3} . Within the measurement error, no deviation in film stoichiometry, crystallinity or density was found for the plasma conditions studied. When a plasma exposure of 1 s was used, the refractive index at 633 nm was 1.37. Despite the fact that LiF is well-known to be transparent, some optical absorption was found in the films. Prolongation of the plasma exposure time leads to an increase in optical absorption while the stoichiometry is unchanged. We therefore consider it plausible to attribute the presence of the absorption features to the interaction of the SF_6 plasma with the film. Also lower growth per cycle values were obtained for longer plasma exposure times, and therefore short plasma exposure times are preferred. It is expected that the optical properties can be further optimized by modifying the plasma conditions. To gain more insight into the deposition process, also QMS measurements were performed. From the detected species it could be concluded the $\text{LiN}(\text{SiMe}_3)_2$ ligand already partly dissociates during the precursor dosing step. In contrast to thermal ALD, additional formation of different fluoromethylsilane and fluorocarbon species was observed during the SF_6 plasma exposure step. Most likely the SF_6 plasma species promote the abstraction of the remaining ligands and their subsequent dissociation in the plasma. Overall, this work demonstrates that the easy-to-handle and readily-available SF_6 plasma is a promising alternative to coreactants such as TiF_4 , or HF-pyridine for ALD of lithium fluorides.



Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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