Structure and dynamics of the fast lithium ion conductor “Li$_7$La$_3$Zr$_2$O$_{12}$”

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The solid lithium-ion electrolyte “Li$_7$La$_3$Zr$_2$O$_{12}$” (LLZO) with a garnet-type structure has been prepared in the cubic and tetragonal modification following conventional ceramic syntheses routes. Without aluminium doping tetragonal LLZO was obtained, which shows a two orders of magnitude lower room temperature conductivity than the cubic modification. Small concentrations of Al in the order of 1 wt% were sufficient to stabilize the cubic phase, which is known as a fast lithium-ion conductor. The structure and ion dynamics of Al-doped cubic LLZO were studied by impedance spectroscopy, dc conductivity measurements, $^6$Li and $^7$Li NMR, XRD, neutron powder diffraction, and TEM precession electron diffraction. From the results we conclude that aluminium is incorporated in the garnet lattice on the tetrahedral 24$\text{d}$ Li site, thus stabilizing the cubic LLZO modification. Simulations based on diffraction data show that even at the low temperature of 4 K the Li ions are blurred over various crystallographic sites. This strong Li ion disorder in cubic Al-stabilized LLZO contributes to the high conductivity observed. The Li jump rates and the activation energy probed by NMR are in very good agreement with the transport parameters obtained from electrical conductivity measurements. The activation energy $E_a$ characterizing long-range ion transport in the Al-stabilized cubic LLZO amounts to 0.34 eV. Total electric conductivities determined by ac impedance and a four point dc technique also agree very well and range from $1 \times 10^{-4}$ to $4 \times 10^{-4}$ S cm$^{-1}$ depending on the Al content of the samples. The room temperature conductivity of Al-free tetragonal LLZO is about two orders of magnitude lower ($2 \times 10^{-6}$ S cm$^{-1}$, $E_a = 0.49$ eV activation energy). The electronic partial conductivity of cubic LLZO was measured using the Hebb-Wagner polarization technique. The electronic transference number $t_e$ is of the order of $10^{-7}$. Thus, cubic LLZO is an almost exclusive lithium ion conductor at ambient temperature.

Introduction

Today, state-of-the-art electrolytes in lithium (ion) batteries are composed of organic solvents or polymers with a dissolved Li-salt. These are combustible in case of abuse, and they often react with electrode materials to form passivating films in the best case. This is on the one hand necessary and favourable, on the other hand proceeding film growth and electrolyte decomposition at elevated temperatures lead to degradation of the battery and shorter battery life times. In particular, new cell concepts based on high voltage cathodes or mobile non-metal cathode systems based on sulfur or oxygen require improved electrolytes, and it may well be that ion-conducting membranes become indispensable. Therefore, the search for stable and incombustible inorganic solid electrolytes with high lithium ion conductivity is currently an important issue in battery research.
Recently Li ion conducting garnet-type oxides are considered as promising electrolytes because of their high conductivity and stability vs. Li metal. Thangadurai et al. first reported about Li ion mobility in Li$_x$La$_3$M$_2$O$_{12}$ ($M = \text{Nb, Ta}$).\(^5\) After the report on cubic Li$_7$La$_3$Zr$_2$O$_{12}$ (LLZO) with the so far highest room temperature conductivity\(^6\) in 2007 these materials have attracted a lot of interest, but it was often found to be difficult to obtain highly conductive LLZO. Later Awaka et al. showed that the reason for this is the existence of a second, tetragonally distorted modification\(^7\) with a much lower conductivity than cubic LLZO. Although some groups reported about the successful preparation of cubic LLZO it is impossible to judge about the quality of the reported materials, due to missing experimental data like conductivities and/or X-ray diffractograms (XRD).\(^8\)\(^9\) Recently, some groups reported on aluminium impurities and the formation of cubic LLZO in combination with Al impurities or Al doping.\(^10\)\(^\text{–}^1\text{3}\) None of these groups could clarify the exact effect of Al on the formation of cubic LLZO and the changes of the crystal structure caused by Al incorporation except the suggestion that Al might substitute Li in the garnet lattice. In order to indicate that it is in fact not pure Li$_7$La$_3$Zr$_2$O$_{12}$ which shows high Li ion conductivity, we will use the abbreviation LLZO or the formula in quotation marks throughout the paper.

Despite attempts to clarify the crystal structure of cubic LLZO by single crystal XRD\(^8\)\(^,^1\text{0}\)\(^\text{–}^1\text{3}\) not much is known about the lithium distribution in cubic LLZO, owing to the small scattering factors of lithium ions. It is the aim of this study to gain deeper insight into the structure and ion dynamics of cubic LLZO in order to better judge its potential as a battery electrolyte. In particular, we made a systematic study of the effect of aluminium doping on the formation of the tetragonal and the cubic LLZO modification. In order to obtain maximal structural information we combined XRD, neutron powder diffraction as well as transmission electron microscopy/precession electron diffraction (TEM-PED) and nuclear magnetic resonance (NMR) spectroscopy. We propose a structure model which takes the lithium ions properly into account as well as the aluminium dopant. In order to also obtain comprehensive information on the lithium ion dynamics we combined electrochemical measurements and "Li NMR spin–spin as well as spin–lattice relaxation techniques. Very good agreement of the different results was found. Finally, we used the Hebb–Wagner method to determine the electronic transference number of cubic LLZO which has not been reported before so far.

**Phase characterization**

Phase analysis along synthesis was carried out on a Siemens D500 Siemens AG, now Bruker AXS in Bragg-Bretano geometry with CuKα-radiation. Phase analysis and determination of cell parameters at room temperature in cubic LLZO were carried out using X-ray powder diffraction (XPD) with a STOE STADI P diffractometer either by using MoKα$_2$-radiation, $\lambda = 0.7093$ Å or CoKα$_2$-radiation, $\lambda = 1.78897$ Å.

Elastic coherent neutron scattering experiments were performed on the high-resolution powder diffractometer SPODI at the research reactor FRM-II (Garching, Germany).\(^1\text{4}\) Monochromatic neutrons ($\lambda = 1.5482$ Å) were obtained at a 155° take-off angle using the (551) reflection of a vertically-focused composite Ge monochromator. The vertical position-sensitive multidetector (300 mm effective height) consisting of 80 $^3$He tubes and covering an angular range of 160°. 2θ was used for data collection. Measurements were performed in Debye–Scherer geometry. The powder sample was filled into a thin-wall (0.15 mm) vanadium can of 13 mm in diameter and then mounted in the top-loading closed-cycle refrigerator. Helium 4 was used as a heat transmitter. The instantaneous temperature was measured using two thin film resistance cryogenic temperature sensors Cernox and controlled by a temperature controller from LakeShore. Two-dimensional powder diffraction data were collected at 4 K and 300 K using two wavelengths and then corrected for geometrical aberrations and curvature of Debye–Scherer rings.

**Experimental**

**Synthesis**

LLZO was prepared by a high temperature route using LiOH (Chempur, 99%, anhydrous), ZrO$_2$ (Chempur, 99.9%) and La$_2$O$_3$ (Chempur 99.99%, dried at 900 °C for 12 h). The starting materials were mixed in stoichiometric amounts (10 wt%–20 wt% LiOH excess was used) and milled for 8 h–12 h in 2-propanol in a planetary ball mill (Fritsch, Pulverisette 7 premium line or Fritsch, Pulverisette 5 classic line) with zirconia balls and grinding bowls. We made use of three different procedures to prepare Al-free and Al-containing LLZO samples. Interestingly, although we followed the preparation route described in the literature\(^6\) to synthesize cubic LLZO, the second calcination step at 1130 °C led to the formation of the tetragonal modification\(^7\) of LLZO. The successful preparation of cubic LLZO turned out to depend on the number of calcination steps and the kind of crucibles used. The cubic phase of LLZO only formed when alumina crucibles were used and when the powder was annealed several times at 1130 °C for about 12 h with milling steps in between. As a result of the reaction between the crucible and the powder a thinning of the crucible after some preparation steps was observed. Several batches of cubic LLZO were prepared according to this procedure and investigated by ICP-OES (inductively coupled plasma optical emission spectroscopy). All of the cubic LLZO samples prepared showed an Al content of about 0.9 wt% (28 mol% Al per mol Li$_7$La$_3$Zr$_2$O$_{12}$). To verify the hypothesis that Al stabilizes the cubic modification we prepared LLZO samples intentionally doped with Al as well as free of Al. For this purpose crucibles free of any Al traces have been used. A small amount (corresponding to 0.5 wt% and 0.9 wt% in LLZO) of γ-Al$_2$O$_3$ (Merck, anhydrous) was added to the mixture of the starting materials. The Al-doped samples were otherwise prepared according to the procedure described above. After calcination at 900 °C and 1130 °C the powder was pressed to pellets, covered with mother powder, and sintered at 1230 °C for 30 h. For comparison, Al-free LLZO has been prepared without addition of γ-Al$_2$O$_3$ powder.
The analysis of powder diffraction data was carried out using the Rietveld technique implemented into the software package FullProf.\textsuperscript{15} The peak profile was described by a modified pseudo-Voigt function (Thompson–Cox–Hastings). The background of the diffraction patterns was fitted using a linear interpolation between selected data points in non-overlapping regions. Cylindrical absorption correction with \( \mu R = 0.7072 \) has been applied to obtain neutron intensities. The scale factor, lattice parameter, fractional coordinates of atoms and their iso-anisotropic displacement parameters, zero angular shift, profile shape parameters and half width (Caglioti) parameters were varied during the fitting. Prior to the Rietveld refinement the lattice parameters, background and profile shape parameters were estimated using the full profile decomposition (Le-Bail) technique.

The maximum entropy method (MEM) implemented in the program PRIMA\textsuperscript{16} was used to determine the nuclear densities. MEM deals with the 3D densities in the way giving the maximum variance of structure factors \( F_o \) within standard deviation of observed structure factors \( F_o \), where structural information can effectively be extracted from the diffraction data and reflected on the resulting three-dimensional distribution of electron/nuclear densities. MEM analysis was conducted using either 189 (at 4 K) or 192 (at 300 K) structure factors obtained from the Rietveld analysis, with the unit cell divided into 128 \( \times \) 128 \( \times \) 128 pixels.

**TEM investigations**

Transmission electron microscopy (TEM) was performed in a F30 STwin microscope (Cs = 1.2 mm, 300 kV, FEG) equipped with a Spinning Star precession module (Nanomegas) and an EDX detector (Fa. Noran). Energy dispersive X-ray (EDX) analyses were performed in a Tecnai F30 STwin microscope (Cs = 1.15 mm, 300 kV, LaB\textsubscript{6}) equipped with a Si/Li detector (EDAX system). The analyses were carried out in scanning transmission electron microscope (STEM) mode using the HAADF detector for image acquisition. The samples for TEM investigations were grinded and suspended in n-butanol. One drop of each suspension was placed on a TEM support (a lacey carbon film on a copper grid). All images were recorded with a Gatan Multi-1 k CCD camera (1 k \( \times \) 1 k) and evaluated with a program Digital Micrograph (Gatan, Inc.). Identification of the crystal modification of LLZO was performed by comparing the diffracted intensities of simulated and experimental precession electron diffraction (PED) images. The precession angle was set to 3\(^\circ\) for the PED experiments. The selected area electron diffraction (SAED) and PED patterns were recorded along different zone axes. Several zone axes were examined, however, in the present work we are focusing on [001] since this direction represents the most significant one for phase distinction. The software JEMS\textsuperscript{17} was used for the simulations of SAED and PED patterns (including multiple scattering events (dynamical calculations)) as well as for simulations of high-resolution TEM (HRTEM) images (using the multislice approach). The Weickenmeier–Kohl atomic form factor was used for the simulations. For PED calculations the sample thickness was set to 10 nm. For HRTEM simulations the following microscope parameters were used: Cs = 1.15 mm, defocus spread 7 nm and illumination seminagle of 0.8 mrad. In order to reduce noise, all HRTEM images were filtered after Fast Fourier transformation using a band-pass mask, with no loss of essential structural information in the following. The structural data for PED simulations of tetragonal LLZO were taken from ref. 7.

**NMR measurements**

\(^7\)Li (spin-quantum number \( I = 3/2 \)) NMR spin–spin- and spin–lattice relaxation rates in both the laboratory and rotating frame of reference were recorded using a modified MSL 100 spectrometer and an MSL 400 spectrometer operating at 4.7 T (77.8 MHz) and 9.4 T (155.5 MHz), respectively. The spectrometers were connected to Oxford cryomagnets. Standard broadband probes (Bruker) as well as home-built high temperature probes were used for data acquisition. 90\(^\circ\) pulse lengths ranged from 4 \( \mu\)s to 6 \( \mu\)s. The temperature in the sample chamber was monitored with the help of an Oxford ITC4 using Ni–CrNi thermocouples to within \( \pm2 \) K. Temperatures below room temperature were adjusted with the help of a flow of freshly evaporated nitrogen. Above 290 K the temperature in the sample chamber was controlled with a stream of heated air. \(^7\)Li NMR spin–lattice relaxation rates (1/\( T_1 \) = \( R_1 \)) in the laboratory frame of reference were recorded using a saturation recovery pulse sequence: a 90\(^\circ\) detection pulse is sent after a comb of closely spaced saturation pulses of the same duration. The corresponding transients \( M_x(t) \) describing the recovery of longitudinal magnetization follow single exponential time behaviour. This is in contrast to the analogous measurements of 1/\( T_1 \) = \( R_1 \) in the rotating frame of reference performed using the spin-lock technique at frequencies of 11.5 kHz and 33.1 kHz. The transients \( M_{xy}(t) \) can be well parameterized by stretched exponentials \( M_{xy}(t) \propto \exp(-t/T_1)^{\gamma} \) with 0.4 < \( \gamma < 1 \).

High-resolution \(^6\)Li and \(^{27}\)Al NMR spectra under fast sample rotation were recorded using an Avance III spectrometer (Bruker) in combination with a shimmed cryomagnet of 14.1 T. Magic angle spinning (MAS) spectra were recorded with a 2.5-mm-probe at 88.4 MHz and 156.5 MHz, respectively. The spinning speed was 30 kHz.

**Electrochemical characterization**

Pellets of sintered LLZO samples were cut into pieces of different lengths and vacuum dried at 100 °C for at least 6 h. The dried pellets were then exclusively handled in an argon filled glove box (MBraun, Labmaster) with less than 0.1 ppm H\textsubscript{2}O and O\textsubscript{2} in the gas. Lithium foil (Chemetall) was used as electrode material for conductivity measurements by two electrode ac impedance spectroscopy and four electrode dc techniques. The lithium foil was pressed on the sides of the LLZO pellets (Fig. 1a) and then annealed at 170 °C for some minutes on a heating plate to improve the contact between Li and LLZO. For the dc measurements two additional electrodes were placed in the flank of the pellet. Therefore, two 0.5 mm wide grooves were cut in the flank and filled with Li metal (Fig. 1b). For both ac and dc measurements the Li
electrodes were contacted with a silver or copper wire. These cells were sealed in gas-tight pouches of aluminized compound foil. Temperature dependent conductivity measurements were carried out in a temperature chamber (WTL 64 Weiss Technik) in the temperature range from $-40 \degree C$ to $100 \degree C$. A potentiostat/galvanostat (model SP-300 from Biologic Science Instruments) was employed for electrochemical characterization. The software package EC-Lab V10.02 was used for data acquisition and impedance data fitting. Impedance spectroscopy was carried out in the frequency range from 7 MHz to 1 Hz (amplitude of 20 mV).

Additionally, low-temperature impedance spectra down to $-148 \degree C$ were recorded by employing a Novocontrol Concept 80 broadband dielectric spectrometer which is equipped with a BDS 1200 sample cell and a BETA analyzer. The latter is capable to measure impedances down to $10^{-14} \Omega$ at frequencies ranging from a few μHz to 20 MHz. Temperature regulation and controlling within an accuracy of about 0.5 K was carried out with a Quattro cryosystem (Novocontrol) using dry nitrogen gas. The root mean square ac voltage was typically 0.1 V to 1.0 V. These data were mainly used to extract electrical relaxation rates from the corresponding modulus spectra.

The dc conductivity was measured by applying a constant current in the range from $1 \mu A$ to $100 \mu A$ between the electrodes 1 and 4 (Fig. 1b) and measuring the resulting voltage drop over the distance of the potential probes 2 and 3 (Fig. 1b). At each temperature a constant dc current was applied for 90 s. The voltage drop was simultaneously measured and remained constant over the measured time period.

The electronic partial conductivity of cubic LLZO was measured using the Hebb–Wagner technique. A gold blocking electrode of approximately 1 μm thickness was deposited on one of the pellets surfaces by thermal Au evaporation. As mentioned above, Li foil was attached on the opposing surface as a reversible electrode for Li ions (Fig. 1c). Measurements were performed in potentiostatic mode (Keithley 6430) in a home-built cell in the range from 2.5 V and 4.5 V vs. Li/Li$^+$ in an argon filled glove box with the Li electrode connected as cathode. Steady-state was assumed when the current remained constant within 10% over a period of several hours.

### Results and discussion

#### Synthesis

Synthesis of cubic LLZO was only successful when an aluminium source was present during synthesis. Thus cubic LLZO formed either by working in Al$_2$O$_3$ crucibles, where the LLZO starting materials react with the crucible, or when intentionally using γ-Al$_2$O$_3$ powder as a dopant in the LLZO synthesis. Preparation of cubic LLZO in Al$_2$O$_3$ crucibles without further doping takes more than two calcination and milling steps due to the stepwise reaction of the powder at the crucible walls only. When the formation of phase pure cubic LLZO (by reaction with the wall) was completed the aluminium content of these samples was typically about 0.9 wt% as determined by ICP-OES. The Al free synthesis did not lead to pure cubic LLZO but rather to the tetragonal modification or a mixture of the cubic and tetragonal modifications of LLZO (Fig. 2a-b). Synthesis with 0.5 wt% Al$_2$O$_3$ led mainly to cubic LLZO, however, the corresponding reflexes are somewhat broader indicating that there might be some tetragonal LLZO present as well (Fig. 2c). Doping with 0.9 wt% Al$_2$O$_3$ in Al free crucibles as well as synthesis without doping but using Al$_2$O$_3$ crucibles led to cubic LLZO (Fig. 2d and e). These results clearly prove that the aluminium content plays a crucial role for the formation of phase pure cubic LLZO.

#### Structural information from MAS NMR

Structural features of Al-doped cubic LLZO were elucidated by $^{27}$Al MAS NMR spectroscopy. In Fig. 3 the NMR spectra

Fig. 2 XRD patterns of LLZO after the final calcination step (a) simulated pattern (JCPDS 80-0457), (b) Al-free synthesis in Al free crucible, (c) doping with 0.5 wt% γ-Al$_2$O$_3$ in Al free crucible, (d) doping with 0.9 wt% γ-Al$_2$O$_3$ in Al free crucible, (e) undoped sample prepared in Al$_2$O$_3$ crucible (0.9 wt% Al content measured by ICP-OES).

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of cubic LLZO before and after sintering at elevated temperatures (see above) are shown. The spectrum obtained after the sintering step is similar to that in ref. 10. Geiger et al. suggest that the signal showing up at a chemical shift of \( \delta_{\text{iso}} = 12 \) ppm reflects octahedrally coordinated Al in LaAlO\(_3\) being an impurity phase.\(^{15}\) The \( \delta_{\text{iso}} \) value of 64 ppm (see Fig. 3b) is characteristic of Al ions residing on lattice sites with tetrahedral coordination. Most probably the Al ions occupy the Li site 24d in the garnet structure according to the aliovalent doping mechanism \( 3\text{Li}^+ \rightarrow \text{Al}^{3+} \). Multiple-quantum magic angle spinning (MQ-MAS) measurements of Geiger et al. have shown that the asymmetry parameter \( \eta \) of the corresponding electric field gradient EFG tensor is close to zero indicating axial symmetry.\(^{10}\) Preliminary \( ^{27}\text{Al} \) MAS measurements on samples with varying Li, La and Al contents indicate that the signal near 80 ppm in fact represents at least two NMR lines.\(^{20}\) Obviously, at a sufficiently large Al content, Al\(^{3+}\) ions occupy various magnetically inequivalent sites in the cubic LLZO structure. NMR lines near 80 ppm, which most likely reflect Al ions also occupying sites with tetrahedral coordination, show up for samples of cubic LLZO which are characterized either by a deficit of La and/or Zr.\(^{20}\)

It is worth mentioning that the cubic LLZO sample obtained before the sintering step (see the \( ^{27}\text{Al} \) MAS NMR spectrum in Fig. 3a) is characterized by a Li ion conductivity which is by about two orders of magnitude lower than that reported for cubic LLZO. The low Li conductivity of the non-sintered sample, which is comparable to that of LLZO crystallizing in tetragonal symmetry, has been confirmed by \( ^{7}\text{Li} \) NMR spin–lattice relaxation measurements revealing a rather low Li diffusivity.\(^{21}\) Comparison of the two spectra reveals distinct differences which might help explain the high ion conductivity (see below) of the sintered material from an atomic-scale point of view. Firstly, the sintering step clearly enhances the Al content within LLZO. Secondly, the NMR line intensity at 12 ppm has decreased after the sintering step. Thus, LaAlO\(_3\) seems to act as a kind of precursor for the incorporation of Al into LLZO.

Finally, in Fig. 4 (top) the \( ^{6}\text{Li} \) MAS spectrum of Al-doped cubic LLZO is shown. Even under MAS conditions with a spinning rate of 30 kHz, the \( ^{6}\text{Li} \) NMR line is comparably broad showing a line width (full width at half maximum, fwhm) of 1.1 ppm. This value should be compared with that one of the tetragonal Al-free counterparts (space group \( I\text{a}d/d\)) (see Fig. 4, bottom).

**Crystal structure from neutron and X-ray diffraction**

Our XRD results do not indicate any tetragonal distortion, but show some weak reflections, which can be attributed to a residue of lithium carbonate. These reflexes are more pronounced in the neutron data (Fig. 5) and the content of lithium carbonate can be quantified as 1.8(4) wt\%. The comparison of neutron powder diffraction patterns collected at 4 K and 300 K indicates the isostructurality of LLZO at both temperatures. All reflexes originate either from cubic LLZO as the main phase or lithium carbonate as the residual phase. Different structure models were considered and simulated\(^{1,10,22}\) which, however, did not reveal a satisfactory description of the Bragg intensities mainly due to underestimated isotropic displacement parameters and differences in lithium occupations. After the adjustment of the models, the best results have been obtained with the model proposed by Geiger et al.\(^{10}\) Aluminium was not taken into account in their structure model, as its position in the lattice cannot easily be determined by X-ray diffraction. In fact, in the presence of relatively heavy lanthanum and zirconium atoms the location of “light” lithium and aluminium atoms is cumbersome. Moreover, the neutron scattering length \( b \) for aluminium is rather low and therefore small concentrations can be hardly quantified using neutron powder diffraction. However, the \( ^{27}\text{Al} \) MAS NMR spectra show that about 70% of the aluminium is located on the lithium sites with 24d symmetry which was included into our structure simulation. Placing Al on the 24d position did not improve the residual intensities in the difference between observed and calculated diffraction patterns, but affects the lithium occupation on the 24d site. Without aluminium \( g_{\text{Li}(24d)} \) has been found to be 42(2)\%, whilst \( g_{\text{Li}(24d)} \approx 52(2)\% \) was...
determined assuming the presence of Al with $g_{Al(24d)} = 6.53\%$. In the latter case the overall lithium content $x$ in Li$_x$La$_3$Zr$_2$O$_{12}$ has been determined to be $6.05 \pm 0.25$, which agrees with the proposed heterovalent substitution mechanism $3\text{Li}^+ \leftrightarrow \text{Al}^{3+}$ (see above).

The best fits to the neutron diffraction data collected at 4 K and 300 K were obtained with the parameters listed in Table 1 and the results of the Rietveld refinements are shown in Fig. 5. Anomalously large isotropic displacement parameters have been noticed for both lithium sites in LLZO already at low temperatures, which might be attributed to static lithium disorder. An application of the split site model to Li on the 24$d$ atomic site did neither lead to a stable fit nor a significant improvement of the fit residuals. The comparison of results obtained at 4 K and 300 K indicates a minor influence of the temperature on the crystal structure of LLZO which is quite typical for garnet-like structures. Heating from 4 K to 300 K results in the elongation of the lattice parameter $a$ by 0.223% with very small atomic movements involved. This means that only a small shift of Li2 in the $a$ direction of the cubic lattice takes place, whereas the coordinates for the other atoms in LLZO remain almost unchanged within the limits of their estimated standard deviations. Temperature increase causes a pronounced rise of displacement parameters on all atomic sites, but the major increase (ca. 50%) was observed for Li on the 96$h$ atomic site and (ca. 30%) for Li on 24$d$.

A strong increase of the displacement parameter indicates a development of dynamic lithium disorder on the background of the static one. The analysis of the temperature evolution of the anisotropic displacement parameters for lanthanum, zirconium and oxygen indicates their evolution from elliptic towards spherical shape with nearly unchanged spatial orientation. This might correspond to a temperature stability of the framework built by these elements. A closer look on the disorder details by examination of differential Fourier maps indicates the presence of quite broad minima around Li (24$d$) with a stripe-like shape penetrating the garnet lattice. This anomaly can be related to the observed very high displacement parameters. However, details of such stripes with negative densities were sufficiently smeared out due to termination effects. Therefore, further examinations of the disorder have been performed using the maximum entropy method (MEM), which can estimate non-zero structure factors for high-Q reflections (typically excluded in the analysis of the powder diffraction data). This feature makes the termination effect in MEM analysis less pronounced in comparison to Fourier synthesis, thus yielding less noisy three-dimensional distributions of electron/nuclear densities. Therefore, the MEM technique has been successfully used to a diversity of scientific problems either concerning disorder in Sr$_{9.3}$Ni$_{1.2}$(PO$_4$)$_7$ and in Rb$_6$Cu$_{11}$(F$_2$Cl$_2$)$_{1.5}$ or experimental visualization of lithium diffusion in Li$_x$FePO$_4$ and La$_{0.69}$(Si$_{0.70}$Mg$_{0.30})$O$_{25.30}$.

The negative nuclear densities obtained after the MEM analysis of datasets collected at 4 K and 300 K are shown in Fig. 6. They correspond to stripes observed in differential Fourier maps and might be related to the lithium diffusion pathway (as lithium is the only negative neutron scatterer in LLZO), where all lithium atoms take part in the diffusion process. Rise of the temperature from 4 K to 300 K unambiguously indicates an enhancement of Li-ion diffusivity, which corresponds to the increase of the ionic conductivity.

**Fig. 5** (a) Experimental and (c) simulated PED patterns of LLZO along [001] assuming the cubic structural model from Table 1. (e) Simulated PED pattern of cubic LLZO according to ref. 9. (g) Simulated PED images of cubic LLZO using the structural model from ref. 10. (b), (d), (f) and (h) Enlarged areas of the marked regions in (a), (c), (e) and (g). The differences between the simulated and experimental PED patterns are clearly seen, with the experimental pattern most convincingly agreeing with the simulated image in (c) (our structure model from Table 1).
It is noteworthy that the structural investigations were carried out on the LLZO powder before sintering. Although the $^{27}$Al NMR results show changes of the Al distribution on different crystallographic sites during sintering which seems to have a significant effect on the Li diffusivity we do not believe that this significantly changes the results of the diffraction investigations. As already mentioned it is quite cumbersome to refine Al and Li positions with a high level of reliability. Therefore, from a crystallographic point of view, the small structural changes detected by NMR after sintering might only lead to an increase of the Li displacement parameters. The small changes in the occupations of La and Zr positions by Al will hardly be detectable by diffraction methods. As the disorder of Li atoms is already quite high, an additional
increase will be within the limit of reasonably refineable displacement parameters. Nevertheless, this will be a subject for further investigations.

**PED and HRTEM results**

In the last decade, PED has become a powerful technique for the advanced characterization of materials. The superior advantage of the PED method over the SAED technique is its ability to reduce dynamical effects arising due to multiple scattering in the transmission electron microscope. Dynamical effects in SAED patterns make the distinction between two phases difficult, particularly, if the phases are structurally and chemically related, as it is the case for cubic and tetragonal LLZO (see ESI†, Fig. SX1). In this case, the diffraction patterns differ only by the absolute values of diffracted intensity. Recently, PED was successfully applied for the distinction of polymorphs occurring in bulk materials and thin films, e.g. of quartz\(^{31}\) and Ag\(_2\)Se\(^{32}\) respectively. In the present work, we compare our results from PED examinations on single microcrystals with those from neutron powder diffraction. Fig. 7a displays the experimental PED pattern of a LLZO sample containing 0.9 wt% of Al and the simulated PED patterns based on the cubic and tetragonal polymorphs also indicate differences in the intensities of (440) and (660) reflections, cf. line profiles in Fig. 7. Thus, the experimental PED pattern is better described by the cubic structure of LLZO. It should be noted that we have tested different structural models for PED simulations obtained from neutron powder diffraction. We found that calculated PED patterns using the structural model in Table 1 show the best match with our experimental PED data. In addition, the calculated PED patterns using the structural model for LLZO of Table 1 match even better as the PED images calculated based on the models for the cubic LLZO\(^{8,10}\) (see e.g. Fig. 8).

Even when adjusting low dose settings, the LLZO sample was extremely unstable under the imaging conditions used for HRTEM. Thus, a complete series of images with variable focus values could not be recorded without initiating considerable loss of structural ordering. Fig. 9a and b show experimental and Fourier-filtered HRTEM micrographs for the first overfocused image of a focus series, zone axis [111]. The simulated micrograph of Fig. 9c convincingly agrees confirming again our assignment to the cubic structure given in Table 1. However, after a few seconds the structure is completely amorphized, just leaving crystalline islands embedded within an amorphous matrix, cf. complete focus series in ESI†, Fig. SX2.

EDX investigations showed a homogeneous distribution of Al in the LLZO sample within the experimental errors. Particularly, no Al-rich regions were found in the LLZO specimen (see ESI†, Fig. SX3).

**Electrochemical results**

Electrochemical measurements were performed to determine the ionic and electronic conductivity of the cubic LLZO. The total conductivity was measured both by ac impedance spectroscopy and four electrode dc measurements. The electronic conductivity was determined by the Hebb–Wagner technique.\(^{18,19}\) Impedance measurements with Au electrodes on LLZO known from the literature show two semicircles and a low frequency tail. These were interpreted to be corresponding to the bulk, grain boundary and electrode contributions, respectively.\(^{6}\) Our impedance measurements performed with
Li electrodes show only one semicircle for the total resistivity of the LLZO pellet and a second semicircle instead of the tail. The appearance of an additional semicircle when using Li electrodes can be expected since Li electrodes can reversibly incorporate Li ions from the LLZO, so the charge transfer resistance and the electrode capacity should give rise to an additional semicircle in the impedance measurements explaining the appearance of a second semicircle instead of the low frequency tail. Still it was surprising that our samples only showed one semicircle for the LLZO contribution so we took a detailed look into the impedance data to be sure that there was no misinterpretation. Fig. 10a shows an example of a typical Nyquist plot measured on a LLZO pellet between Li electrodes. Impedance data were fitted with an equivalent circuit (RC)(RQ) or (RQ)(RQ) depending on whether the respective sample showed a depressed high frequency semicircle or not.

Fig. 8 (a) Experimental and (b) simulated PED patterns of LLZO along [001]. For the simulation of (b) we used the cubic structure model presented in Table 1. (c) Simulated PED images of tetragonal LLZO based on the structure model from ref. 7. (d)–(f) Line profiles taken from (440) and (660) reflections in (a)–(c). The differences in the intensities of (600), (440) and (660) reflections are clearly seen. The experimental pattern fits better to the cubic LLZO.

Fig. 9 (a) Experimental HRTEM micrograph of LLZO along [111] viewing direction. (b) Fourier-filtered HRTEM image of the region marked in (a). (c) Simulated HRTEM image (defocus 90 nm, $t = 9$ nm, 2-fold astigmatism estimated to 10 nm) based on the crystal structure model of LLZO from Table 1 (inserted).
From the fit to this measurement the first high frequency semicircle resistance and capacity were determined to be 20.4 kΩ and 0.38 nF, respectively. The second low frequency semicircle resistance and capacity result in 13.3 kΩ and 0.15 μF, respectively. These relatively high capacity values are typical for grain boundaries or electrodes of polycrystalline ceramics. As both semicircles yield to comparable resistances as supposed for the bulk and grain boundary contributions in cubic LLZO, some additional experiments were carried out to clarify the origin of the impedance contributions when using Li electrodes. Fig. 10b shows a Nyquist plot of a sample measured once with Au and once with reversible Li electrodes. The Au electrodes were gas phase deposited by thermal evaporation, whereas the Li electrodes were pressed mechanically on the pellet. The measurement with the Au electrodes shows a semicircle merging into a low frequency tail. The low frequency tail is caused by an almost solely capacitive behaviour of the Au electrodes indicating their blocking nature under the given experimental conditions. The measurement with Li electrodes leads to two semicircles proving that the second semicircle corresponds to the Li electrodes charge transfer resistance and capacity. This also clarifies that our samples only show one semicircle representing the total resistivity of the LLZO pellet. Murugan et al. reported that bulk and grain boundary contributions could only be resolved at low temperatures, i.e., in their case room temperature and slightly above. In the present case even at −40 °C only one semicircle representing the total resistivity of the sample is observed. As shown in Fig. 10c the charge transfer resistance of the Li electrodes depends strongly on the preparation method of the Li electrodes. Two impedance measurements of the same sample, in one case with Li electrodes gently pressed on the pellet and in the second case with Li electrodes fused on the pellet, were performed. The different preparation methods change the charge transfer resistance almost by one order of magnitude while the pellet resistance remains unchanged as seen in the inset. We did not find any evidence for a reaction of the LLZO with Li metal which might have shown up as an additional impedance during our measurements. It has to be noted that the charge transfer resistance is not negligible and may play a role when LLZO is applied as a solid electrolyte in battery cells with Li metal electrodes. The total voltage drop across a cell is the sum of the charge transfer resistances at the electrode interfaces and the voltage drop along the electrolyte bulk itself. Therefore further investigations on the charge transfer resistances between different electrode materials and preparation procedures are in progress.

In addition to ac impedance we used four electrode dc measurements to determine the conductivity of the samples under continuous current load. By using the four electrode dc technique the bulk resistance of the sample (as a sum of grain and grain boundary contributions) can be determined without electrode overvoltages. The results of the dc conductivities are shown together with ac data in Fig. 11. Comparing the conductivities of samples with different aluminium contents it is clearly visible that the conductivity depends on the aluminium content in the samples. The highest conductivity of $3 \times 10^{-4}$ S cm$^{-1}$ to $4 \times 10^{-4}$ S cm$^{-1}$ at 25 °C were found for samples containing 0.9 wt% Al prepared either by synthesis in Al$_2$O$_3$ crucibles or by intentional γ-Al$_2$O$_3$ doping. The Arrhenius activation energy is determined to be $0.34$ eV. The conductivity and activation energy data agree well with results from Murugan et al. who did not report on any aluminium in their samples and did not give information on the impurity level at all. The 0.5 wt% Al containing sample (γ-Al$_2$O$_3$ doped) shows a slightly lower conductivity of $1 \times 10^{-4}$ S cm$^{-1}$ but the same activation energy of $E_a = 0.34$ eV. The Al-free sample with tetragonal crystal structure is characterized by a two orders of magnitude lower conductivity of $2 \times 10^{-6}$ S cm$^{-1}$ at 25 °C and an activation energy $E_a = 0.49$ eV. The latter value is in good agreement with that reported for a tetragonal sample which has recently been

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**Fig. 10** Room temperature impedance spectra of different LLZO samples with various electrodes (a) impedance spectrum of a LLZO pellet doped with 0.9 wt% Al. The pellet size was 5.3 mm diameter and 16.2 mm length. The pellet was sandwiched between two Li electrodes (frequency range from 7 MHz to 50 mHz at 20 mV amplitude). An equivalent circuit (RC)(RQ) was used to fit the impedance data. (b) Impedance spectra of a LLZO pellet doped with 0.9 wt% Al measured with blocking Au and reversible Li electrodes in a frequency range 7 MHz to 1 Hz at 20 mV. (c) Impedance spectra of a LLZO pellet doped with 0.9 wt% Al measured with Li electrodes pressed and fused on the pellet (frequency range 7 MHz to 1 Hz at 20 mV).
investigated by NMR spectroscopy. \(^{21}\) The electronic conductivity of cubic LLZO, which controls the self-discharge of a lithium battery and therefore is of great interest, was measured with a Hebb–Wagner type cell arrangement with one reversible and one blocking electrode (Fig. 1c). In optimal cases Hebb–Wagner measurements can be used to determine the individual electron hole and electron partial conductivities as a function of the Fermi level (i.e. as a function of the lithium metal content in the present case).

When the cell is polarized, as can be seen in Fig. 1c, a high mixed ionic and electronic transient current is rapidly decreasing and finally reaching a steady state electronic current. The steady state current under ion-blocking conditions can only be due to electron and hole conduction in the material as no Li ions can be delivered from the blocking gold electrode. Prior to the polarization experiments the open circuit voltage of the cell shown in Fig. 1c was approximately 2 V which is probably due to formation of a lithium gold alloy at the LLZO Au interface\(^ {34}\) prior to the measurement. Lithium activity gradients in the sample may also be a reason as well as capacitive effects of the high resistance cell. (Note that the low electronic conductivity leads to an electronic resistance in the order of \(10^{10}\) Ω). Fig. 12 shows the potential dependent mean conductivities measured in the range of typical battery operating voltage. As described in detail in the literature on the Wagner–Hebb technique, the increasing voltage leads to a corresponding decrease of the chemical potential of Li at the Au electrode. This changes the electron and hole concentrations in LLZO. Therefore, the overall electronic partial conductivity increases with increasing polarization voltages. Over the potential range of 2.5 V–4.5 V the electronic conductivity takes values in the range from 5 \(\times\) \(10^{-12}\) Scm\(^{-1}\) to 5 \(\times\) \(10^{-11}\) Scm\(^{-1}\). With the total conductivity of 3 \(\times\) \(10^{-4}\) Scm\(^{-1}\) the electronic and ionic transference numbers

\[
t_{el} = \frac{\sigma_{el}}{\sigma_{total}} \quad \text{and} \quad t_{li^+} = \frac{\sigma_{li^+}}{\sigma_{total}}
\]

result in values between 2 \(\times\) \(10^{-8}\)–2 \(\times\) \(10^{-7}\) and 0.99999998–0.9999998, respectively. This clearly shows that cubic Al-doped LLZO is an almost exclusive Li ion conducting material having great potential as an electrolyte for lithium-ion batteries. We note that the increase of the electronic conductivity with increasing potential is due to growing hole conduction.

**Solid-state NMR measurements**

In addition to impedance spectroscopy we used various NMR techniques to probe Li dynamics in a sintered sample of garnet-like LLZO crystallizing with cubic symmetry. In Fig. 13 the central transitions (see below) of a typical set of \(^{7}\)Li NMR spectra of cubic LLZO doped with 0.9 wt\% Al are shown. The corresponding \(^{27}\)Al MAS NMR spectrum is shown in Fig. 3b. The \(^{7}\)Li NMR spectra were recorded at various temperatures at a resonance frequency of 155.5 MHz. The broad NMR line, which is observed at the very low temperatures, can be well represented by a Gaussian reflecting the distribution of resonance frequencies due to non-averaged dipole–dipole interactions of the Li nuclei. Here, the associated line width \(\delta\) (full width at half maximum) of this so-called rigid lattice regime is given by \(\delta_{rl} = 9.7(2)\) kHz. In this temperature range the mean Li jump rate \(1/\tau\) is much smaller than the spectral width \(\delta_{rl}\), i.e., \(1/\tau < 10^3\) s\(^{-1}\). With rising temperature \(1/\tau\) increases resulting in an averaging of homonuclear dipolar interactions. Consequently, the NMR line increasingly narrows, finally reaching the width \(\delta_{mn}\) which is solely governed by inhomogeneities of the external magnetic field used (regime of extreme narrowing). Interestingly, in the case of cubic LLZO doped with Al a heterogeneous motional narrowing (MN) is observed. At intermediate temperatures...
the NMR line partially narrows resulting in a two-component line shape, i.e., the broad rigid-lattice line (see above) is superimposed by a motionally narrowed Lorentzian-shaped component. Recently, the same behaviour has been observed by Koch et al. who studied the Li dynamics in the garnet Li$_5$La$_3$Nb$_2$O$_{12}$. It is worth mentioning that the spectra shown in Fig. 13 were simply obtained by Fourier transformation of a free induction decay recorded after excitation with a 90° pulse. Thus, quadrupole intensities arising from the interaction of the spin-3/2 probe nucleus with electric field gradients produced by the electric charge distribution in the neighbourhood of the Li nuclei are largely suppressed. These can be made visible by using appropriate echo pulse sequences. Usually, when low temperatures are regarded, such interactions are represented by a broad Gaussian line of low intensity characterized by a width of some tens of kHz. As an example, a stimulated echo NMR spectrum is shown in Fig. 13e which has been recorded using the Jeener–Broekaert echo sequence. In the present case, we will exclusively focus on the $^7$Li NMR central transition of LLZO. In contrast to other garnet-like oxides, in the case of cubic LLZO two-phase NMR spectra show up at temperatures as low as 200 K indicating a spin ensemble of rapidly diffusing Li ions (see Fig. 13). As an example, at 220 K the narrow Lorentzian-shaped component shows a line width of approximately 1.9 kHz while the broad one is characterized by $\delta = 8.7$ kHz. The area fraction $A_f$ of the narrow line, which reflects the number fraction of fast Li ions with jump rates larger than about $10^4$ s$^{-1}$, amounts to approximately 19% at this temperature. Values of $A_f$ have been estimated by fitting the spectra with a combination of two Voigt functions. Interestingly, $A_f$ steadily increases with increasing $T$ reaching 28% at 233 K, 35% at 243 K, and 41% at 253 K. At a temperature of 263 K nearly 50% of the total number of Li ions participate in a very fast Li diffusion process. The corresponding line width of the sharp component became smaller than 1 kHz. Thus, with rising temperature the fractional amount of Li ions being highly mobile on the NMR time scale increase in an almost linear fashion (Fig. 13c), a similar behaviour has been found for nanocrystalline LiNbO$_3$. Let us note that at higher temperatures a separation of the two spin reservoirs with their distinct dynamics turns out to be difficult. This is due to the fact that the broad spectral component, representing rather
slow Li ions in LLZO, steadily decreases in intensity. The heterogeneous MN observed at low \( T \) might reflect a large distribution of Li jump rates. Such a distribution is expected to be preserved even at temperatures larger than 300 K. Interestingly, even the line width of the sharp component reveals a stepwise narrowing reflecting a transition from partial to full averaging of dipolar interactions. Starting from 2 kHz at 263 K the line width is further decreased with increasing \( T \) finally reaching 800 Hz in the regime of extreme narrowing (see Fig. 13d). This is very similar to the behaviour observed by \(^7\)Li NMR spin–spin-relaxation measurements discussed below (see Fig. 14).

In addition to NMR line shapes, \(^7\)Li NMR spin–lattice relaxation (SLR) rates have been recorded in both, the laboratory and rotating frames of reference, see ref. 21, 37, and 39 for an introduction into these NMR techniques. It is worth mentioning that the two components of the NMR line are characterized by the same \(^7\)Li SLR NMR rates \( R_1 \) and \( R_\text{pp} \). Thus, a separation of the two, dynamically different spin reservoirs according to that shown previously for the nanocrystalline two-phase Li conductor Li_xO:Al_{2}O_3,35,40 is not possible. In contrast to the present work, in the previous study two spin ensembles, which seem to be unaffected by spin-diffusion effects, were anticipated to be also spatially separated.29 The temperature dependence of the rates of the sintered sample of cubic LLZO, which have been recorded at different frequencies, are shown in Fig. 14 in an Arrhenius plot. Below 180 K, i.e., in the rigid-lattice regime (see above), the rates are governed by non-diffusive background relaxation and reveal a weaker-than-activated temperature dependence. Presumably, the SLR NMR rates in this temperature range are influenced by lattice vibrations, interactions between the spins and paramagnetic impurities and/or by strictly localized, i.e., caged ion dynamics which do not reflect translational jump processes. However, at higher temperatures the SLR NMR rates pass through characteristic diffusion-induced rate peaks from which Li jump rates \( 1/\tau \) and activation energies \( E_{a,NMR} \) can be deduced. Interestingly, the rates recorded at 77.8 MHz reveal an asymmetric diffusion-induced peak \( R_1(1/\tau) \). In general, the low-temperature flank, for which \( \omega_0 \tau \gg 1 \) holds, is influenced by short-range Li motions which might be affected by correlation effects (see below).21,37,39,40

Compared with this, on the high-temperature side of the rate peak, i.e., in the limit \( \omega_0 \tau \ll 1 \), the SLR rate is governed by long-range Li transport.21,37,39,40

The mean diffusion parameters characterizing the latter are directly comparable with the results obtained from dc conductivity measurements (see above). The solid line in Fig. 14 represents a fit according to the model introduced by Bloembergen, Purcell and Pound41 which was originally developed for isotropic uncorrelated diffusion: \( R_1 \propto \tau / (1 + \omega_0 \tau)^b \).

Here, the asymmetry of the rate peak is taken into account by the parameter \( \beta \), which can adopt values ranging from 1 to 2. \( \beta = 2 \) is the so-called BPP-behaviour, is obtained for uncorrelated motion. \( \beta < 2 \) indicates that the underlying hopping correlation function \( G(t) \) is represented by a non-exponential rather than an exponential function. Correlation effects such as structural disorder and/or Coulomb interactions are considered to be responsible for the deviation from exponential time behaviour of \( G(t) \). In the present case, \( \beta \) amounts to about 1.4; the activation energy \( E_{\alpha,NMR} \) turns out to be approximately 0.31 eV. This value is in good agreement with that probed by impedance spectroscopy (see above). For comparison with \( E_{\alpha,NMR} \), the corresponding activation energy in the limit \( \omega_0 \tau \gg 1 \) turns out to be about \( E_{\alpha,NMR} = 0.12 \) eV. In general, \( E_{\alpha,NMR} \) and \( E_{\alpha,NMR} \) are linked with each other via \( E_{\alpha,NMR} = (\beta - 1) E_{\alpha,NMR} \).37 In addition, \( \beta \) determines the frequency dependence of \( R_1 \) in the limit \( \omega_0 \tau \gg 1 \) according to \( R_1 \propto \omega_0^b \). No frequency dependence is expected in the limit \( \omega_0 \tau \ll 1 \) when 3D diffusion is regarded.37 This is in contrast to ion conductors showing low-dimensional Li diffusion.37,42–44

Owing to technical limitations of our NMR setup, temperatures higher than 800 K could not be reached. Thus, the high-temperature flank of the \( R_1 \) rate peak recorded at 77.8 MHz was only partly accessible. However, with the help of SLR NMR measurements performed in the rotating frame of reference and carried out at locking frequencies \( \omega_0 \approx 2 \pi \) in the kHz the complete high-temperature flank of the corresponding diffusion-induced rate peak can be probed. The dotted line in Fig. 14 represents an Arrhenius fit of some of the data points of the high-\( T \) flank of the \( R_{1p} \) data recorded at \( \omega_0 \approx 2 \pi \approx 11.5 \) kHz. The fit yields an activation energy of approximately 0.34 eV and is in good agreement with the trend of the \(^7\)Li NMR spin–spin relaxation rates \( R_2 \) which are also included in Fig. 13. The \( R_{1p} \) measurements are limited by \( R_2 \) leading to an apparently broad rate peak whose top is cut off. Nevertheless the temperature at which the rate maximum appears can be estimated to be approximately 300 K. In general, the diffusion-induced SLR rate maximum shows up when the Li correlation rate \( 1/\tau' \) reaches the order of the angular frequencies \( \omega_0 \) and \( \omega_1 \), respectively.37 \( 1/\tau' \) is expected to be identical with the mean jump rate \( 1/\tau \) within a factor of two. In Fig. 15 the Li

![Fig. 14](Image)

**Fig. 14** Temperature dependence of the \(^7\)Li NMR spin–spin \( (R_2) \) as well as spin–lattice relaxation rates \( (R_1 \) and \( R_{1p} \) of cubic LLZO. The NMR spin–lattice relaxation rates were measured at the frequencies indicated. Spin–spin relaxation rates were recorded at a resonance frequency of 77.8 MHz. See text for further details.
jump rates $1/\tau_{\text{NMR}}$ directly deduced from the rate maxima according to $\omega_{0}^{2} = \frac{1}{\tau_{\text{NMR}}} \approx 0$, see ref. 39, respectively, are shown in an Arrhenius plot. The additional vertical axis in Fig. 15 converts the rates $1/\tau_{\text{NMR}}$ into self-diffusion coefficients according to the Einstein–Smoluchowski equation: $D_{\text{NMR}} = \frac{a_{j}^{2}}{6\tau_{\text{NMR}}}$. The mean jump distance $a_{j}$ was estimated to be approximately 0.2 nm. In analogy to previous studies, comparing results obtained from NMR with those from impedance spectroscopy, dc conductivity values $\sigma_{\text{DC}}$ (Fig. 11) have been roughly converted into Li jump rates using both the Nernst–Einstein and Einstein–Smoluchowski equations. As is well-known, the combination of the two relations yields

$$1/\tau_{\text{NMR}} = \frac{6k_{B}T}{Nq^{2}a_{j}^{3}}\sigma_{\text{DC}}$$

where $N$ denotes the number density of charge carriers and $q$ the charge of the Li ions. The product of Haven ratio $H_{R}$ and correlation factor $f_{c}$ was assumed to be 1. Additionally, in Fig. 15 electrical relaxation rates $1/\tau_{M}$ are included which were obtained from the imaginary part of the electrical modulus $M''$ which has been analyzed as a function of frequency $\nu$. Modulus spectra $M''(\nu)$ were recorded at various temperatures $T$ (not shown here for the sake of brevity). The characteristic maxima show up when $1/\tau_{M''}$ is of the order of the angular frequency $\nu 2\pi$. The solid line in Fig. 15 represents a fit taking into account $1/\tau_{M''}$ and $1/\tau_{c}$ yielding 0.34 eV (see above).

The jump rates $1/\tau_{\text{NMR}}$ obtained from NMR $R_{1}$ measurements are in fair agreement with the conductivity results. The agreement between $1/\tau_{c}$ and $1/\tau_{\text{NMR}}$ probed via $R_{\text{DC}}$ turns out to be somewhat better which might be due to the fact that the two methods probe Li dynamics on a very similar time scale. For the sake of completeness, a Li jump rate is included in Fig. 15 which has been estimated from NMR line narrowing (Fig. 8). Slightly above 200 K the narrow component shows up. At 225 K the corresponding line width is reduced to $\delta_{0}^{1/2}$ and the associated jump rate is given by $1/\tau_{\text{NMR}} = \delta_{0}^{1/2}$. For comparison, at 205 K the rates $R_{2}$ start to deviate from their rigid-lattice value $R_{2D} = 60 \mu$s leading to $1/\tau_{\text{NMR}} \approx R_{2D}$ at this temperature. This jump rate is also included in Fig. 15. Taken together, the jump rates probed by NMR spectroscopy corroborate the transport parameters of a sintered sample of cubic LLZO obtained from impedance spectroscopy. In agreement with line widths measurements (Fig. 13d), $R_{2}$ shows a two-step decrease with increasing temperature indicating that the Li ions get access to further diffusion pathways at elevated temperatures.

**Conclusions**

We have synthesized the two different structural modifications of LLZO and found a certain Al content to be necessary for the formation of cubic LLZO. One already could speculate from the patent by Yoshida\textsuperscript{47} that Al is a necessary ingredient for the formation of cubic LLZO. As in the present work, the recently published papers\textsuperscript{10–13} state that cubic LLZO only forms when Al is present during synthesis but there was no definite knowledge about the role of Al. With the combination of XRD, TEM-PED and neutron scattering we are able to propose a structure model which takes both Al and Li ions into account. Neutron diffraction measurements show a high degree of Li disorder even at 4 K which becomes even more pronounced at 300 K. We regard this disorder as the reason for the high mobility of the Li ions in the cubic garnet-type lattice. As proved by $^{27}$Al MAS NMR spectroscopy most of the Al ions are located on the 24$d$ site. With the results of phase transformation during synthesis we regard the Al ions on the 24$d$ site to stabilize the cubic garnet structure.

NMR was found to be a very sensitive probe for small structural changes in the garnet-type lattice. NMR shows that minor changes of the Al distribution during the sintering process cause the Li disorder to become even more pronounced than in the powder before sintering. The TEM-PED patterns agree very well with our proposed structure model and the corresponding pattern simulation; other structure models fit less well. The simulated structure image also fits well with the HRTEM image, confirming our cubic structure model.

Electrochemical measurements performed with different LLZO samples confirm that the conductivity depends on the Al content, i.e., the degree of transformation from the tetragonal to the cubic crystal structure. Ac and dc conductivities show almost identical values approving the high conductivity also under current load. Li jump rates deduced from NMR spin-lattice relaxation measurements agree well with those calculated from electrical conductivity data. Moreover, the activation energies determined by both techniques agree very well. The electronic conductivity and the corresponding electronic transference number are very small, and LLZO was found to be stable against reaction with Li metal over the measured period of several days. Therefore, we suppose that Al-stabilized cubic LLZO is a suitable Li solid electrolyte. Tests of the compatibility of LLZO with various cathode materials and measurements of the overvoltages occurring
in contact with these as well as Li metal under current load are in progress and will help further clarify the applicability of cubic LLZO as a solid state electrolyte for Li ion batteries.

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Notes and references