INTRODUCTION

The direct conversion of heat into electricity by thermoelectric generators is a promising approach to generate power for sensor systems in hot and harsh environments. For this special application, thermoelectric oxides are especially interesting, as they can be used at high temperatures up to 800°C in air. Thermoelectric materials in general are evaluated by the power factor $PF = S^2 \cdot \sigma$ and the figure of merit $ZT = (PF \cdot T)/\kappa$. For infinite heat sources as in the field of energy harvesting, the thermoelectric material should be optimized for maximum $PF$ to enable a high-power generation.1 For a limited heat source as in the field of energy recovery, a high-energy conversion efficiency is required and thus a maximum $ZT$ is important.1 Thermoelectric oxides, although having lower $ZT$ than conventional thermoelectric materials like tellurides and skutterudites,2 are an interesting alternative at elevated temperatures as they are stable in air. In

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Abstract

Calcium cobaltite $\text{Ca}_3\text{Co}_4\text{O}_9$, abbreviated Co349, is a promising thermoelectric material for high-temperature applications in air. Its anisotropic properties can be assigned to polycrystalline parts by texturing. Tape casting and pressure-assisted sintering (PAS) are a possible future way for a cost-effective mass-production of thermoelectric generators. This study examines the influence of pressure and dwell time during PAS at 900°C of tape-cast Co349 on texture and thermoelectric properties. Tape casting aligns lentoid Co349. PAS results in a textured Co349 microstructure with the thermoelectrically favorable ab-direction perpendicular to the pressing direction. By pressure variation during sintering, the microstructure of Co349 can be tailored either toward a maximum figure of merit as required for energy harvesting or toward a maximum power factor as required for energy harvesting. Moderate pressure of 2.5 MPa results in 25% porosity and a textured microstructure with a figure of merit of 0.13 at 700°C, two times higher than the dry-pressed, pressureless-sintered reference. A pressure of 7.5 MPa leads to 94% density and a high power factor of 326 µW/mK² at 800°C, which is 11 times higher than the dry-pressed reference (30 MPa) from the same powder.

KEYWORDS

hot pressing, texture, thermoelectric properties

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1 | INTRODUCTION

The direct conversion of heat into electricity by thermoelectric generators is a promising approach to generate power for sensor systems in hot and harsh environments. For this special application, thermoelectric oxides are especially interesting, as they can be used at high temperatures up to 800°C in air. Thermoelectric materials in general are evaluated by the power factor $PF = S^2 \cdot \sigma$ and the figure of merit $ZT = (PF \cdot T)/\kappa$. For infinite heat sources as in the field of energy harvesting, the thermoelectric material should be optimized for maximum $PF$ to enable a high-power generation.1 For a limited heat source as in the field of energy recovery, a high-energy conversion efficiency is required and thus a maximum $ZT$ is important.1 Thermoelectric oxides, although having lower $ZT$ than conventional thermoelectric materials like tellurides and skutterudites,2 are an interesting alternative at elevated temperatures as they are stable in air. In
addition, their raw materials are less toxic and more abundant. One of the most promising oxide p-type materials is calcium cobaltite $\text{Ca}_3\text{Co}_4\text{O}_9$ (in the following abbreviated by $\text{Co349}$). It has a layered misfit crystal structure of alternating layers of a hexagonal $\text{CoO}_2$ sublattice and a triple rocksalt type $\text{Ca}_3\text{Co}_4\text{O}_9$ sublattice. $\text{Co349}$ has a plate-like, lentoid grain morphology due to an increased grain growth in ab-direction. Due to its crystal structure, the physical and thermoelectric properties of single-crystal $\text{Co349}$ are strongly anisotropic. The electrical conductivity of single-crystal whiskers is 500 to 1000 times higher in ab-direction than in c-direction ($\sigma_{ab} = 500\ldots1000\times\sigma_c$). Hot-pressed samples with 10 MPa show anisotropic Seebeck coefficients $S_{ab} = 1.06 \times S_c$ and anisotropic thermal conductivity $\kappa_{ab} = 2.04 \times \kappa_c$. $\text{Co349}$ decomposes at 926°C. This limits the sintering temperature and thereby the densification in the case of conventional sintering.

The anisotropy of the particle can be assigned to the polycrystalline material by texturing the microstructure. A detailed literature review about the texturing of $\text{Co349}$ can be found in our previous work or by Prasad et al. Here we only focus on some aspects of texturing $\text{Co349}$. Dry-pressing with high pressure up to 1000 MPa is a simple way to increase texture, but the sample size is limited. Casting a slurry composed of anisotropic particles and organic binders by the doctor-blade method into a tape is a well-known technique to texture materials. The effect can be enforced by implementing a fraction of large anisotropic grains ($>10 \mu m$), known as templated grain growth. The tape-cast specimens by Schulz et al. showed anisotropic microstructure and high porosity (45%). Due to the latter, electrical conductivity and $PF$ were not improved compared with the dry-pressed reference (300 MPa). Nagahamo et al. doubled the electrical conductivity using templated grain growth and tape casting $(600 \text{ S/m at 400°C})$ compared to a nontextured compound fabricated by cold isostatic pressing. Applied pressure during sintering increases both sinter density and texture. Several groups applied hot-pressing (or sinter-forging, respectively) with pressures from 5 MPa to 50 MPa. The $\text{Co349}$ specimens were highly textured and nearly fully dense. With a pressure of 30 MPa, Kenfaui et al. reached $\sigma(900K) = 8 \times \sigma_{\text{ref}}$. Due to the enhanced charge carrier mobility in ab-direction, the thermal conductivities of hot-pressed specimens measured in ab-direction are enhanced by the factor of 10 or 4.7 compared to the reference. Liu et al. and Delorme et al. used spark plasma sintering (SPS) with a pressure of 50 MPa ($70 \text{ MPa}^2$). A relative density of $> 98\%$ was reached with $\sigma = 12 000 \text{ S/m} = 3.5 \times \sigma_{\text{ref}}$. Comparing SPS and hot-pressing, the latter leads to a higher anisotropy and higher $PF$. Finally, a combination of texturing during shaping at room temperature and texturing during sintering is possible. Schulz et al. combined tape casting and pressure-assisted sintering. Pressure-assisted sintering is a sintering technology known from the field of low-temperature co-fired ceramics (LTCC) using a moderate uniaxial pressure during sintering and a different pressing plate setup than hot-pressing. Pressures of 7.5 MPa and 10 MPa during sintering led to nearly fully dense specimens with very high electrical conductivities in ab-direction ($\sigma(1000K) = 17 000 \text{ S/m}^2$).

Tape casting has much potential for texturing $\text{Co349}$ as shown by Schulz et al. It is widely used in the field of electroceramics, for example, for LTCC, piezo actuators, or lambda-probes, and it can be used to design thermoelectric generators (TEG). It is easily scalable and an approach for inexpensive mass production of TEG-stacks. Due to the low density of conventionally sintered $\text{Co349}$, the produced parts are fragile. Based on the work of Schulz et al., we recently reported on the combination of tape casting and pressure-assisted sintering to texture $\text{Co349}$. In this work, we focused on the influence of powder composition on the thermoelectric properties when using pressure-assisted sintering. We found that Bi-doping ($\text{Ca}_{2.7}\text{Bi}_{0.3}\text{Co}_4\text{O}_9$) is not beneficial as it leads to distortions and abnormal grain growth due to a liquid phase when a pressure of 10 MPa is applied during sintering. The microstructure of the samples from undoped $\text{Co349}$ fabricated with pressure-assisted sintering showed strong texture, high sinter density, and thus both electrical conductivity and $PF$ in ab-direction were strongly increased by more than a factor of 10 compared to the dry-pressed sample from the same powder. Due to the high density ($94\%$), $\kappa_{ab}$ increased as well for the PAS-samples, which nearly compensated the benefits of texturing regarding $ZT_{ab}$. It would be favorable to apply a moderate pressure during PAS to benefit from texturing on the one hand and to maintain a certain porosity to decrease $\kappa_{ab}$. Hence, in this work we focus on the influence of pressure and dwell time during PAS on the thermoelectric properties of undoped $\text{Co349}$ to optimize either $PF$ or $ZT$.

## Experimental Procedure

$\text{Ca}_3\text{Co}_4\text{O}_9$ was synthesized by solid-state reaction. Stoichiometric amounts of the raw material $\text{Co}_3\text{O}_4$ (99.97% ChemPUR, Karlsruhe, Germany) and $\text{CaCO}_3$ (99%, low-alkali, Riedel-de Haën, Seelze, Germany) were attrition-milled (moliNEx, Netzsch, Selb, Germany) for 45 minutes at 800 rpm using $\text{ZrO}_2$ grinding media ($d = 2.5 \text{ mm}$). The powder mixture was calcined at 900°C for 12 hours in air to obtain $\text{Ca}_3\text{Co}_4\text{O}_9$. The calcined powder was ground in a planetary ball mill (Pulversette 5, Fritsch, Idar-Oberstein, Germany) for 20 minutes with agate grinding balls and subsequently attrition-milled for 15 minutes to obtain $\text{Ca}_3\text{Co}_4\text{O}_9$ powder with a $d_{50} = 3 \mu m$ (particle size distribution determined with laser diffraction system, Mastersizer 2000, Malvern Instruments, Malvern, United
Kingdom) and a specific surface of 3 m²/g (BET, NOVA 2200, 3P INSTRUMENTS, Odelzhausen, Germany). This powder was mixed with polyvinyl butyral (Eastman, Kingsport, US), dibutylphthalate (ACROS organics, Fair Lawn, US), rhodafac RE-610 (Solvay, Hannover, Germany), ethanol (Merck, Darmstadt, Germany), methylethylketone (Merck, Darmstadt, Germany) and cyclohexanone (Merck, Darmstadt, Germany) in a porcelain mill for one day to obtain a tape-casting slurry. The slurry was tape-cast with doctor-blades on a tape-casting machine (Netsch, Selb, Germany) to produce a ceramic green tape with a mean thickness of 70 µm and an organic content of 12 wt% (loss on ignition at 450°C). The tape was cut into sheets with a length of 50 mm and sintered at 900°C for 24 hours in air. Reference samples were prepared by dry-pressing of Co349 particles. Not to scale

FIGURE 1 Scheme of the simplified manufacturing process of the PAS laminates. Small black rectangles illustrate the orientation of Co349 particles. Not to scale.
and the thermoelectric voltage $\Delta U$ was measured in-between the platinum contacts of the thermocouple. A modulated temperature gradient $\Delta T$ of around 20 K over the bar-like specimens was applied by an additional heater. The contacted samples were mounted in a special sample holder placed in a gas-flushable tube furnace, which was purged by dry synthetic air. The measurements were performed stepwise from 900°C to room temperature. The Seebeck coefficient $S$ was calculated as described in reference 32. The measurement setup is indirectly validated with another Seebeck measurement system.

Minitab 17 was used to statistically analyze the data. The data samples were tested for differences in mean value with two-sample t-test and for differences in standard deviation with F-test with a level of significance set to 5%. The sample size (number) of the room temperature data was at least 5, and of the high-temperature data at least 3.

### 3 | RESULTS AND DISCUSSION

#### 3.1 | Phase content, microstructure, and texture

The XRD patterns of the starting Co349 powder, the sintered test bars, and the sintered laminates with different pressure and 2 h dwell time; reference values for Ca$_3$Co$_4$O$_9$ taken from 8 [Color figure can be viewed at wileyonlinelibrary.com]

![XRD patterns](image)

| $p$ in MPa | $t$ in h | rel. Density in % | MRD | $S$ in $\mu$VK$^{-1}$ | $\sigma$ in Scm$^{-1}$ | $PF$ in $\mu$Wm$^{-1}$K$^{-2}$ | $\kappa$ in Wm$^{-1}$K$^{-1}$ | $ZT$ in 10$^{-3}$ |
|-----------|----------|------------------|-----|----------------------|------------------|----------------------|----------------------|----------------------|
| 0 (ref)   | 2        | 49.9 ± 0.2       | 3.5$^9$ | 120.2 ± 0.2       | 9 ± 1             | 14 ± 6               | 0.40 ± 0.01           | 12 ± 2               |
| 0         | 2        | 57.2 ± 0.1       | 9.2  | 124.0 ± 0.5        | 29 ± 5            | 44 ± 1               |                       |                      |
| 2.5       | 2        | 74.8 ± 4.4       | 12.1 | 124.6 ± 0.3        | 94 ± 31           | 146 ± 48             | 1.70 ± 0.01           | 26 ± 9               |
| 2.5       | 4        | 78.3 ± 1.3       | 123.9 ± 0.3 | 96 ± 14       | 148 ± 21           |                       |                      |
| 2.5       | 8        | 80.7 ± 0.3       | 124.5 ± 0.4 | 126 ± 28      | 195 ± 43           |                       |                      |
| 5.0       | 2        | 84.3 ± 1.5       | 126.1 ± 0.5 | 128 ± 29      | 203 ± 45           |                       |                      |
| 7.5       | 2        | 93.5 ± 0.8       | 135.4 ± 6.1 | 151 ± 11      | 279 ± 43           | 4.03 ± 0.02           | 21 ± 3               |
| 7.5       | 8        | 94.7 ± 0.3       | 146.9 ± 0.5 | 140 ± 2       | 302 ± 6            |                       |                      |

Note: Relative density, $S$, $\sigma$, and $\kappa$ (all at room temperature) are given with standard deviation. For $PF$ and $ZT$, the error according to Gaussian error propagation is given. MRD of dry-pressed test bar is derived from reference 9.
pattern as the calcined powder, the intensity ratio between (111) and (004) reflection is 0.16 smaller than the value of the powder (0.37). This indicates a weak texturing due to dry-pressing. The (001) reflections of the tape cast and pressureless-sintered laminates are strongly increased, supporting the assumption of a texturation during tape casting. The PAS laminates show only (001) peaks, indicating a very strong texture. The laminate sintered with 2.5 MPa shows an additional peak at 22°. This could be the (111) peak of Co$_3$O$_4$. Co$_3$O$_4$ is formed as a reaction layer between the release tape (Al$_2$O$_3$) and Co349. Usually, the reaction layer was removed after sintering. The peak indicates an incomplete removal of the reaction layer.

Figure 3 illustrates the microstructure of the different specimens. The calcined powder (see Figure 3A) has the typical flake-like morphology of Co349. Due to the milling, the single-crystal layers are visible. As shown in Figure 3B, the grains are rounded after sintering at 900°C without pressure (reference sample – not tape-cast). Sinter necks are visible, but the structure is still very porous. Sintering without pressure is not sufficient to obtain a dense microstructure at 900°C. The sinter necks and pores indicate that the thermal treatment only leads to a first stage sintering. After tape casting, a slight particle orientation is visible in the SEM images (see Figure 3C). The pressureless-sintered laminate shown in Figure 3D) has a slightly different microstructure than the sintered test bar. The grains are smaller and the microstructure is still very porous. Considering the results from the XRD measurements (see Figure 2), tape casting leads already to a textured microstructure. As illustrated in Figure 3E) the fracture surface of a laminate sintered with a uniaxial pressure of 5 MPa for 2 hours is very dense without large pores. The laminar crystal structure is visible. It is difficult to detect single grains at a fracture surface or a polished surface. Therefore, the polished surfaces of the PAS specimens were thermally etched at 900°C for 30 min (Figure 3F-H). Etching reveals that the microstructure includes pores. The grains are between 1 and 3 µm long and about 1 µm thick. All thermally etched specimens show fine white particles (l ≈ 160 nm, h ≈ 80 nm). EDX-point analysis (8kV, 150 000×) of nearly free-standing white particles revealed that they are Co-enriched (Co-L: 41 at% versus 29 at% for the matrix) and Ca-deficient (Ca-K: 5 at% versus 16 at% for the matrix). We, therefore, suppose that these are Co$_3$O$_4$-nanoparticles, probably an effect of polishing and thermal etching as these particles are not found in the fractured samples. As shown in Figure 3F-H), the number of pores decreases with increasing pressure. At 7.5 MPa the microstructure appears to be completely dense (see Figure 3H). An increased dwell time does not seem to increase the densification remarkably (Figure 3F-I). Comparing Figure 3F) with Figure 3I), longer dwell times increase the grain size from 1.5 ± 0.7 µm to 2.4 ± 0.8 µm. It can be concluded that pressure-assisted sintering is a good way to obtain a dense textured microstructure of Co349.

Pressure during sintering works as an additional driving force and therefore is an effective way to support densification. Higher pressure apparently leads to improved densification. Longer dwell times increase the grain size but not the densification.
To determine the influence of the pressure during sintering on texture, the latter was analyzed using XRD pole figures of (002) and (111) reflections as shown in Figure 4. All tape cast samples show a strong fiber texture with a preferential orientation of the c-axis orthogonal to the tape casting direction. Multiples of a random distribution (MRD) in out of plane direction were calculated from estimated orientation density function (ODF) data (MTEX Software) to quantify the texturing (see Table 1). The MRD can vary from 1 (no texture) to infinity (single crystal). As the texture of the dry-pressed bar was not measured, in Figure 5A the MRD data of a similar processed Co349 test bar were used as a reference 9. The reference exhibits a low degree of texture. The tape cast sample prior to sintering already shows a strong fiber texture (MRD = 9.3). Sintering without pressure does not increase the texture effect (Figure 4A). PAS increases the texturing with the c-axis parallel to the pressing direction (see Figure 4B-D and Figure 5A). With increasing pressure, the texture gets stronger (Figure 4C) and reaches a maximum MRD of 12.8. Longer dwell times do not increase the texturing as shown in Figure 4D).

3.2 | Density

The relative density (after sintering) of the dry-pressed samples is with 50% very low (Table 1) in comparison with other
But in these studies, higher pressures during dry-pressing (up to 300 MPa) were applied, resulting in higher green-densities and thus higher sinter-densities (60% to 72%). Tape casting increases the relative density to 57%, but the parts are still very fragile when no pressure is applied during sintering. Increased density due to tape casting is described in literature. With increasing uniaxial pressure, the relative density increases to 94% for 7.5 MPa (Figure 5A). The increase in the relative density as a function of pressure during sintering fits a linear regression within this experimental framework. The relative density of the PAS specimens is in the range of the hot-pressed samples observed by Kenfaui et al.9

### 3.3 Seebeck coefficient

The Seebeck coefficient at room temperature $S_{(22°C)}$ increases by 3% when tape casting is used (Figure 5B). With increasing pressure, $S_{(22°C)}$ increases further. The correlation between $S_{(22°C)}$ and pressure is not linear. At a pressure level of 7.5 MPa, $S_{(22°C)}$ has an unusual high standard deviation. According to literature, $S$ is higher in ab-direction than in c-direction (for hot-pressing with 10 MPa, $S_{ab} = 1.06S_c^{9,10}$). Due to the particle orientation during tape casting, $S_{(22°C)}$ is increased in ab-direction. As PAS also leads to a further texturing (see Figure 5A), the increase of $S_{(22°C)}$ with pressure is attributed to enhanced texture. The Seebeck coefficients at 100°C of PAS samples are in the same range as the values measured at room temperature (Figure 6B). $S_{ab}(100°C)$ of the dry-pressed reference is 20 µV/K lower than measured at room temperature. $S_{ab}$ of all three samples increases with temperatures until it reaches a maximum at 800°C. The slope of the specimens sintered with 7.5 MPa (linear fit between 300°C and 700°C, slope 0.04 µV/K$^2$) is less steep than of the two other samples (linear fit between 300°C and 700°C, slope 0.07 µV/K$^2$). There is no significant difference between $S_{ab}$ of the reference and of the PAS sample pressed with 7.5 MPa above 700°C (2-sample t-test). Above 400°C, the specimens sintered with 2.5 MPa show the highest $S_{ab}$ values, with a maximum of $S(800°C) = 164$ µV/K. The literature data for hot-pressed Co$^{349}$ are slightly higher in the range 170-180 µV.9,10,16

### 3.4 Electrical conductivity

The electrical conductivity at room temperature of the dry-pressed specimens is very low$^2$ (Figure 5C). Tape casting increases $\sigma_{(22°C)}$ by the factor of 3. With increasing pressure, $\sigma_{(22°C)}$ increases strictly monotonically. Although, the increase in conductivity from 2.5 MPa to 5 MPa and from 5 MPa to 7.5 MPa are not significant, respectively (two-sample t-test). The large differences between the experimental points in room temperature electrical conductivity remain almost unchanged up to 900°C (Figure 6A). The temperature has only a minor influence on $\sigma_{ab}$. Due to the low density.

![Figure 6](https://example.com/figure6.png)
of the dry-pressed sample, its electrical conductivity is lower as for similar studies.\(^9\),\(^14\) But as shown in Figure 6A, PAS increases the electrical conductivity compared with both data-sets for dry-pressed samples. The results are in the same range as for hot-pressed Co349 from other studies.\(^9\),\(^16\) Kenfaui et al\(^9\) reported 11,500 S/m at room temperature for hot-pressed samples with 5 MPa, Nagahama et al\(^18\) derived 14,000 S/m at 200°C for hot-pressed samples with 8 MPa and Schulz et al\(^16\) found 12,500 S/m at room temperature for hot-pressed samples with 10 MPa. In the latter study, the temperature dependency was much higher with an increase in 35%.

3.5 | Power factor

The dry-pressed reference has a low PF at room temperature. Tape casting increases the PF\((22°C)\) by the factor of 3.1 due to texture. PAS with 2.5 MPa results in an increase by a factor of 10.4, and PAS with 7.5 MPa leads to a 20 times higher PF\((22°C)\) with respect to the reference as both \(\sigma\) and \(S\) are higher (Figure 5). Over temperature, PF increases for all three measured samples (Figure 6C). It reaches a maximum at 800°C. The results are \(PF_{ab}(800°C) = 30\ \mu W/mK^2; 231\ \mu W/mK^2\), and 326 \(\mu W/mK^2\) for reference, 2.5 MPa, and 7.5 MPa, respectively. This corresponds to an increase by a factor of 10.9 or 5 compared to the dry-pressed sample of this study or the one of Kenfaui et al\(^9\), respectively. The slopes of the PAS samples are steeper than of the dry-pressed reference. Kenfaui et al\(^9\) and Schulz et al\(^16\) reached slightly higher PF-values \((T_{max} = 700°C)\), due to the above-explained differences in \(S\) and \(\sigma\).

3.6 | Thermal conductivity and figure of merit

Phonon and charge carrier transport are both higher along the crystal layers of Co349.\(^10\) Thus, the increase in \(\sigma(22°C)\) and density in ab-direction of the textured microstructure is accompanied by an increase in thermal conductivity \(\kappa(22°C)\) (see Table 1) up to a factor of 10. Measuring \(\kappa\) of hot-pressed samples in pressing direction (c-direction) underestimates \(\kappa\) as the phonon transport is interfered by the crystal layers as, for example shown in reference 16. Combining the moderate increases in \(\kappa\) and the strong increase in PF, the highest \(ZT_{ab}(22°C)\) of 0.026 is reached for PAS samples sintered under a pressure of 2.5 MPa. For all measured specimens, \(ZT_{ab}\) increases with temperature (Figure 7). Contrary to PF and \(\sigma\), the sample sintered with 2.5 MPa shows the highest values over the full temperature range. With \(ZT_{ab}(700°C) = 0.13\), it is almost two times higher than \(ZT\) of the reference. The difference between the sample sintered with 7.5 MPa and the reference is not as high as for the other properties due to the much higher thermal conductivity, \(\kappa_{ab}\), of the 7.5 MPa sample. For a maximum \(ZT_{ab}\), it is not beneficial to produce a microstructure as dense as possible. Although high densities enhance \(PF_{ab}\) largely, \(ZT_{ab}\) is reduced due to high \(\kappa\) in this direction. Schulz et al\(^16\) reported very high \(ZT\)-values (0.29 at 600°C) for tape cast Co349 pressureless-sintered samples. As \(\kappa\) was measured in c-direction, the \(ZT\)-value is probably overestimated. The only other \(ZT_{ab}\) for hot-pressed Co349 was published by Kenfaui et al.\(^10\) For hot-pressed samples with 10 MPa, a maximum \(ZT_{ab}(750°C)\) of 0.16 was reached. Kenfaui determined \(\kappa_{ab}\) for the whole temperature range. In literature,\(^10\),\(^16\)

### TABLE 2

Calculated \(ZT\) at 700°C \(ZT(700°C)\) is calculated using \(\kappa(22°C)\) as measured in this study. \(ZT_{fit}(700°C)\) is calculated with \(\kappa_{fit}(700°C)\)

| \(PF(700°C)\) in \(\mu W/mK^2\) | \(\kappa(22°C)\) in \(W/mK\) | \(ZT(700°C)\) with \(\kappa(22°C)\) in \(10^{-3}\) | \(\kappa_{fit}(700°C)\) in \(W/mK\) | \(ZT_{fit}(700°C)\) with \(\kappa_{fit}(700°C)\) in \(10^{-3}\) |
|---|---|---|---|---|
| reference | 30 | 0.44 | 66 | 0.43 | 67 |
| 2.5 MPa, 2 h | 225 | 1.70 | 129 | 1.11 | 197 |
| 7.5 MPa, 2 h | 327 | 4.03 | 79 | 2.23 | 142 |

Note: All bold values were calculated using the extrapolated \(\kappa_{fit}(700°C)\) according to \(\kappa_{fit} = A \cdot e^{-\frac{T}{B}} + C\) with \(A = 0.849, B = 512, C = 0.892\) for 2.5 MPa derived from Shi et al\(^{36}\) and \(A = 2.106, B = 205, C = 2.166\) for 7.5 MPa.\(^{15}\)
κ of hot-pressed Co349 decreases with rising measuring temperature. The κ over temperature curve is very similar for similar κ in different studies\textsuperscript{10,16,35,36} and it can be fitted by the following equation $\kappa = A \cdot e^{-\frac{T}{T_0}} + C$ with absolute temperature $T$, and constants A, B, and C. In this study, only κ of the reference was determined over the whole temperature range. $\kappa_{ab}$ was only measured at room temperature and then used to approximate $ZT_{ab}$ for the whole temperature range. Thus, the $ZT_{ab}$ values presented in this paper are probably underestimated. When $ZT_{ab}$ is calculated using the extrapolated $\kappa_{fit}$ based on the κ over temperature curve of similar κ from literature,\textsuperscript{35,36} the $ZT$ values at 700°C as shown in Table 2 are reached ($ZT_{ab}$). This results in $ZT_{fit}(700°C)$ of 0.2 for 2.5 MPa, which is higher than published for hot-pressed Co349 using $\kappa_{ab}$.\textsuperscript{10,37}

### 3.7 | Influence of dwell time

The results in Table 1 do neither show a significant influence of the dwell time on the relative density nor on the conductivity at room temperature $\sigma(22°C)$ or on $PF(22°C)$ (two-sample $t$-test). Only the Seebeck coefficient $S(22°C)$ at a pressure level of 7.5 MPa is increased significantly by an increased dwell time. It can be concluded that a prolonged dwell time during PAS is not beneficial for the thermoelectric properties.

### 3.8 | Effect of density and texture

The data shown in Figure 5 and Table 1 suggest a dependency of the electrical conductivity and the Seebeck coefficient at room temperature, $\sigma(22°C)$ and $S(22°C)$, respectively, on the apparent density of the specimens. Figure 7 shows the respective plots. The Seebeck coefficient increases non-linearly with density. The reference has the lowest density and the lowest $S(22°C)$. In the range of 2.6 g/cm$^3$ to 4.0 g/cm$^3$, $S(22°C)$ is almost constant. The range comprises the tape cast and the PAS samples sintered with a pressure up to 5 MPa. The samples with a density above 4.3 g/cm$^3$ show by far the highest Seebeck coefficient. The graph leads to the assumption that the increase in $S(22°C)$ for tape cast samples is due to the texture and not due to the increased density, as $S(22°C)$ remains nearly constant when the density is further increased. The almost fully dense specimens have a very high Seebeck coefficient, probably due to a combined mechanism of density and texture. The Seebeck coefficient of Co349 is anisotropic.\textsuperscript{10} A higher density of the sintered test bars leads to higher electrical conductivity, as shown in Figure 7. The correlation can be even fitted with a linear regression ($R^2(adj) = 94\%$). But the linear correlation between electrical conductivity and density is false positive as texture and density increase simultaneously with increasing pressure during sintering. When the data presented in this paper are compared to PAS specimens with a similar density but a lower degree of texture\textsuperscript{12} as derived by reaction sintering (see Table 3), it is obvious that a lower degree of texture (MRD = 4.1) leads to a lower electrical conductivity. Kenfaui et al\textsuperscript{22} showed a similar effect for highly dense Co349 prepared by SPS (low MRD) and hot-pressing (high MRD). This means that the increase in electrical conductivity over pressure (see Figure 5C) is a combined mechanism of increased density and increased texture. A higher density enlarges the intergranular contact areas as porosity is decreased simultaneously. This leads to the observed higher electrical conductivity. Previous studies have shown similar dependencies for oxide thermoelectric materials.\textsuperscript{11,31} A higher degree of texture increases the electrical conductivity in ab-direction due to the anisotropy of the material.\textsuperscript{22}

### 4 | SUMMARY

This study about the influence of dwell time and pressure during pressure-assisted sintering of Co349 elucidates the following points:

1. Tape casting yields textured samples (MRD = 9.2) when conventional sintering is used. Pressure-assisted sintering (PAS) of tape-cast samples slightly increases the texture (MRD ≈ 12) and considerably increases the density.
2. A prolonged dwell time during hot-pressing is not beneficial for the thermoelectric properties. Longer dwell times do not densify the samples but coarsen the microstructure.
3. By pressure variation during sintering, the microstructure of Co349 can be tailored either toward maximum $PF_{ab}$ as required for energy harvesting or toward maximum $ZT_{ab}$ as required for energy recovery. This means lower

| MRD | $\rho$ in g/cm$^3$ | $\rho_{rel}$ in % | $S$ in $\mu$V/K | $\sigma$ in S/m | $\kappa$ in W/mK |
|------|----------------|------------------|------------|------------|-------------|
| Calcined, 7.5 MPa, 2 h | 12.8 | 4.38 ± 0.04 | 94 ± 1 | 135 ± 6 | 15 140 ± 1182 | 4.03 |
| Reaction-sintered, 7.5 MPa, 2 h\textsuperscript{12} | 4.1 | 4.24 ± 0.00 | 91 ± 0 | 142 ± 2 | 9763 ± 1157 | 2.91 |

Note: Data for the reaction-sintered sample are derived from reference\textsuperscript{12}.
pressure for a porous microstructure and maximum $ZT_{ab}$, higher pressure for full densification and maximum $PF_{ab}$.

4. Higher applied pressures during sintering result in higher densities, in higher electrical conductivities $\sigma_{ab}$, in higher Seebeck coefficients $S_{ab}$, and higher power factors $PF_{ab}$. A pressure of 7.5 MPa increases $PF_{ab}(700^\circ C)$ by more than one order of magnitude to 326 $\mu W/mK^2$ compared to the dry-pressed reference from the same powder.

5. Measuring the thermal conductivity $\kappa_{ab}$ for Co349 with the hot-disc method is an appropriate method to be applied for flat specimens to obtain all thermoelectric data in the same direction.

6. With increasing pressure during sintering, $\kappa_{ab}$ increases. A pressure of 7.5 MPa increases $\kappa_{ab}(22^\circ C)$ by the factor of 10 compared to the reference. With respect to $ZT$, the high $\kappa_{ab}$ nearly compensates the increase in $PF$ for high-pressure levels.

7. The highest ZT was measured for PAS samples pressed with a moderate pressure of 2.5 MPa to $ZT_{ab}(700^\circ C) = 0.13$, which is almost the double value compared to the dry-pressed reference. Lower pressures during sintering are a good compromise between texturizing to enhance $\sigma_{ab}$ and relative high porosity (25%) to maintain a low $\kappa_{ab}$.

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**ORCID**

Sophie Bresch  
https://orcid.org/0000-0003-2490-7208

Björn Mieller  
https://orcid.org/0000-0002-0784-9790

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