

Micro- and Macro-Mechanical Testing of Transparent $\mathrm{MgAl_{2}O_{4}}$ Spinel

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ABSTRACT

The mechanical properties and reliability aspects of advanced transparent fineand coarse-grained MgAl₂O₄ spinel have been characterized at ambient and high temperature. The studies were based on a combination of micro- and macro-mechanical methods to assess Young's modulus, hardness, fracture toughness, strength and crack growth kinetics. The results and reliability aspects are discussed in terms of linear elastic fracture mechanics. Strength was analyzed using two- and three-parameter Weibull statistics. The experimental limits of Young's modulus determination using standard ringon-ring testing are highlighted, and an approach is outlined to correct the measured apparent values. Experimentally obtained strength data as a function of loading rate are used to assess the potential effect of subcritical crack growth, yielding the failure time under static loading, via a strength/probability/time plot for a lifetime prediction. Furthermore, the Brinell indentation showed potential for local strength measurement and the data supported a strength-loaded area relationship that was based on the ring-on-ring test data. Novel experimental methods were implemented for fracture determination and strain detection. Particular attention was directed to the effect of defects and associated local strain fields, as assessed using polarized light in photoelastic measurements. Complementary fractography by optical, confocal and scanning electron microscopy provided a correlation between failure initiating defect size and fracture stress.

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KURZFASSUNG

Eigenschaften und Zuverlässigkeitsaspekte fortschrittlicher Mechanische transparenter fein und grobkörniger MgAl₂O₄ Spinelmaterialien wurden bei Raumtemperatur und erhöhten Temperaturen untersucht. Die Untersuchungen basierten auf einer Kombination mikro- und makromechanischer Methoden zur Bestimmung von Young's Modul, Härte, Risszähigkeit, Festigkeit und Risswachstumskinetik. Resultate und Zuverlässigkeit wurden unter Berücksichtigung der linear elastischen Bruchmechanik diskutiert. Die Analyse der Festigkeiten basierte auf Zwei- und Dreiparamter-Weibullstatistiken. Die experimentellen Grenzen der Young's Modul Ermittlung im Fall eines Standard Doppelring-Biegetests werden hervorgehoben und eine Korrekturmethode der scheinbaren experimentellen Werte ist beschrieben. Experimentell als Funktion der Lastrate ermittelte Festigkeitsdaten wurden zur Beurteilung des potenziellen Effekts des unterkritischen Risswachstums verwendet und erlaubten eine Lebensdauervorhersage anhand von Festigkeits - Wahrscheinlichkeits - Zeit Graphen. Weiterhin wurde das Potential der Brinell Härteeindrucksmethode zur Messung der lokalen Festigkeit gezeigt und berechnete Daten stimmten mit einer auf Doppelringmessungen basierten Gleichung des Zusammenhangs Festigkeit - belastete Fläche überein. Neuartige experimentelle Methoden wurden zur Festigkeits- und Dehnungsdetektion eingesetzt. Besondere Aufmerksamkeit wurde auf den Effekt von Defekten und dem damit im Zusammenhang stehenden lokalen Dehnungsfeld gerichtet, wobei Untersuchungen auf photoelastischen Messungen beruhten. Komplementäre fraktographische Untersuchungen mittel optischer, konfokaler und Rasterelektronen -Mikroskopie ergaben einen klaren Zusammenhang zwischen versagensrelevanten Defekten und Bruchspannung.

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1 INTRODUCTION

Ceramics are increasingly utilized in a variety of technical applications, where either specific functional or structural properties (or both) are needed [¹]. In particular, transparent ceramics have gained importance in technical window applications [²]. Spinel is a transparent oxide ceramic currently under consideration for such applications. One of the industrial development targets is to produce highly reliable transparent ceramics at low cost. Magnesia spinel (MgAl₂O₄) appears promising for attaining the envisioned goals by modifying materials and improving the fabrication process. In general, for such functional applications, the mechanical properties must be sufficient to warrant the structural integrity of the component. Therefore, in addition to detailed characterization of the mechanical properties (strength and hardness in particular), analyzing failure origins is a critical aspect.

Results obtained from mechanical measurements and microstructural characterizations in this study are presented for the considered MgAl₂O₄ material. In addition, a main focus of this study is to evaluate different micro- and macro-mechanical characterization methods, particularly the utilization and verification of macroscopically non-destructive characterization approaches, and the correlation of results with microstructural features (i.e., grain size, defects). As a novel experimental approach, an indentation testing device developed in-house and permitted in-situ through thickness fracture observation and photoelastic measurements was used for this transparent material to obtain information on the variation of local stress fields.

Elastic modulus, fracture toughness, and fracture strength as important basic materials properties have been characterized thoroughly. In addition, the effect of environmentally assisted subcritical crack growth and its implication on long-term reliability via a strength/probability/time plot was considered.

Of equal importance to the statistical data for materials development and improvement is the knowledge of failure-relevant defects, particularly the effect of imperfections and associated local stress fields. Here, fractographic analysis based on the combination of confocal and scanning electron microscopy with advanced software packages provides important information.

2 LITERATURE REVIEW

2.1 Spinel materials

The name "spinel" is used for the ternary oxide compound MgAl₂O₄ [³] and also as a name of a group of chemical compounds with an identical crystallographic structure and the general formula – AB_2O_4 , where A is a divalent charged atom such as Mg⁺², Fe⁺², Mn⁺², Zn⁺², and B is a trivalent charged atom such as Al⁺³, Fe⁺³, Cr⁺³, Pb⁺³. Due to its chemical compatibility with alumina, zirconia and mullite, spinel is also considered to be an attractive matrix for ceramic matrix composites [⁴]. Some typical examples of the spinel group are aluminates (MgAl₂O₄), ferrites (MgFe₂O₄), chromites (FeCr₂O₄), franklinites (ZnFe₂O₄), gahnites (ZnAl₂O₄), magnetites (Fe₃O₄) and miniumes (Pb₂PbO₄). The physical and chemical properties of spinel are governed by the A and B compounds as well as by the distribution of cations located at the different crystallographic sites [⁵]. The *fcc* cubic spinel crystal structure (*Fd3m*) is a close-packed array of oxygen ions with the general formula AB_2O_4 , in which Mg⁺² cations occupy 8 of the 64 tetrahedral interstices and Al⁺³ cations occupy 16 of the 32 octahedral interstices [⁶,⁷] (Fig. 2). The lattice parameter A_o of stoichiometric spinel is 0.80832 nm [⁸].

Property optimization motivates continuing efforts to produce new materials, extend phase fields and improve homogeneity [^{9,10,11}]. This, in turn, provides the impulse to develop new synthesis and processing approaches.

The basic procedures to produce spinel materials have been known since 1905 [¹²] and the system MgO·Al₂O₃ was defined in 1916 [¹³] and has essentially remained unchanged since then. Already since the 1930s, attempts have been made to apply magnesium aluminate spinels and use them instead of magnesia chromite products, even though the latter have a lower price and are simpler to manufacture. This is due to an increasing awareness of the toxicity of the Cr⁺⁶ formed by Cr₂O₃ under alkaline conditions, as well as stricter regulations for the use and waste disposals of refractories

containing chromium species [¹⁴]. The awareness of toxicity increased the industrial importance of aluminates (MgAl₂O₄) and ferrites (MgFe₂O₄).

In the 1960's, spinel was already recognized for its high hardness, strength and optical transparency [¹⁵,¹⁶]. Transparent polycrystalline spinel was initially developed as a transparent armor material in 1969 [¹⁷]. It was fabricated using classic ceramic technologies: preparing high purity ultrafine spinel powder, pressing the powder into pellets and finally sintering.

Spinel still did not reach much commercial success until the middle 1980s, mainly due to its complex processing. Many studies exist on the effect of processing parameters on the properties of spinel and its composites, both stoichiometric and non-stoichiometric (magnesia or alumina rich compounds) $[1^{18}]$. At the beginning of the 1980s, polycrystalline spinel was produced by hot-pressing spinel powder, resulting in optical properties equivalent to that of single crystals $[1^{19}]$; starting around the same time, metal oxide spinels have been synthesized by various methods based on the mechanical oxide and/or salt mixtures calcination [20, 21, 22]. Recently, micron-grained and sub-microngrained spinels have been produced by means of extremely homogenous slip-casting with acceptable mechanical properties, as well as high transparency in the visible and infrared spectral range [²³]. Currently, micron-, submicron- and nano-grained spinels ceramics are under development, and already show many processing and property advantages over conventional coarse-grained spinel $[^{24}]$. In particular, many methods have been developed to synthesize high quality powders as ingot material $\begin{bmatrix}25, 26\\ 2, 3\end{bmatrix}$. However, the lack of suitable, well-developed and low-cost preparation methods still limits spinels' application as highperformance transparent ceramics.

2.2 Application

In general, spinel compounds are of interest since they exhibit a wide range of novel and adjustable properties, making the material useable for electronic, magnetic, catalytic, photonic and structural applications [²⁷,²⁸].

In particular, the high melting point of the magnesium aluminate spinel (2135 °C), high mechanical strength at ambient and elevated temperatures, good thermal shock resistance and chemical inertness are an important combination of properties, which has

already led to the application of magnesium aluminate spinels in metallurgy, electrochemistry and chemical industry [29, 30, 31], mainly:

• Refractories

The high melting point and excellent thermal and chemical resistance makes spinel, here as a major component in an alumina-or magnesia-rich matrix, useable for refractory applications. Due to large tensile hoop stresses, the thermal expansion mismatch between periclase and spinel phases in certain areas of phase diagram (20wt.%, see Fig. 1) leads to micro-crack development around the periclase grains in the spinel matrix if a certain grain size is exceeded. This decreases strength and stiffness, but also can reduce the probability of crack propagation in thermal shock loading [³²]. The major application areas of spinel refractories are transition and burning zones of cement rotary kilns, sidewalls and bottoms of steel ladles, because they are resistant to corrosion by slag [³³]. Ordinary ceramic refractory materials typically show a heterogeneous porous microstructure, which consists of large (max. grain size is typically 3 - 5 mm) and fine grains, which form the matrix. Although magnesium aluminate spinels are known to present an attractive combination of physical and mechanical properties, their technological application is, due to the difficulties in sintering and quality issues, still limited to magnesia clinkers used for the construction of furnaces [³⁴].

Optical Devices

Sintered polycrystalline fine-grained spinel with its high refractive index, low scatter and random orientation of small crystallites, is optically completely isotropic. In the visible wavelength range (380 - 750 nm), it exhibits a similar optical transmission as the single crystalline material (> 80 %), with some differences at short ultraviolet (UV) wavelengths (230 - 400 nm). Transparent ceramics have attracted great attention for their use in a wide range of optical applications: high-power lasers, electro-optic devices and lenses. They also have been considered to be an optical material for infrared sensors [35,23]. UV-transparent materials also attract significant interest as laser lenses in the microelectronic chip industry [1]. Markets for portable digital devices have shown strong growth potential, requiring the use of compact lenses, light sensors and protective transparent shields. Frequently, a compact and small lens systems design is necessary for use in cell phones and cameras, and gives rise to the development of new transparent

materials with a high refractive index and high Abbé number, since this enables downsizing of optical systems [³⁶]. In addition to the refractive index, dispersion value is also important for optical components. Particularly for camera lenses, low dispersion materials are preferred in the correction of chromatic aberration [³⁷].

• Armor materials

Transparent armor is a system of functionally-integrated transparent materials whose main role is to provide blast/ballistic protection of windshields and side windows, while retaining the structural integrity and optical transparency of the component [³⁸]. Armor systems are traditionally laminates of soda-lime glass layers. A means to reduce weight and dimensions is the partial replacement of glass by transparent ceramics, since the ceramic has higher hardness, Young's modulus and fracture toughness. Armor performance has not been successfully correlated to a single material property to date, due to the dynamic nature of the ballistic event. However, several fundamental material properties have been used to rank ceramics for screening purposes, with hardness appearing to be the main parameter.

• Energy technology applications

Highly porous magnesium aluminate spinel is considered a suitable inert substrate in some solid oxide fuel cell (SOFC) designs [³⁹]. For employing spinel material as SOFC substrate, the material must be porous, to allow access of gases to the electrodes. Here, high strength (even in the porous state) and chemical inertness are important prerequisites. An additional advantage for this application is that its coefficient of thermal expansion can be varied by changing the amounts of the constituent phases, which limits thermally-induced stresses.

• Other areas where spinel application is considered are: insulating material for fusion reactors and nuclear waste containers; dental implants; tribological applications; gas purging cones; ceramic gas nozzles for welding and plasma cutting; and refractory lances $[1, 2^{8}, 4^{0}]$.

2.3 The MgO·Al₂O₃ System

Spinel (MgAl₂O₄) is a stable intermediate compound in the binary phase diagram $MgO - Al_2O_3$ [⁶]. The limits of the diagram are MgO and Al₂O₃, which have rather high melting temperatures of 2800 °C and 2020 °C, respectively (Fig. 1).

The solid-state reaction between MgO and Al₂O₃ is governed by counter-diffusion of Al⁺³ and Mg⁺² ions through the oxygen lattice of the spinel phase. The diffusion of three Mg⁺² ions is compensated by two Al⁺³ ions in the opposite direction and, therefore, three moles of spinel are formed at the Al₂O₃ - MgAl₂O₄ interface for every mole formed at the MgO-MgAl₂O₄ interface [⁴¹,⁴²]. As a result, if a stoichiometric initial composition of MgO and Al₂O₃ is used by slowly heating and cooling down, at the peritectic point a pure spinel is obtained. In many cases, the compound MgO-*n*Al₂O₃ is non-stoichiometric, i.e., *n* is not equal to 1 and can range from 1 to 7.3, since excess Al⁺³ ions occupy tetrahedral sites, substituting Mg⁺² ions. This causes a proportional decrease of the lattice parameter due to the smaller diameter of Al⁺³ [⁴³]. Pure spinel has a volume expansion of 5 – 7% [⁴⁴] and melts at 2135 °C, with a eutectic point at 1995 °C. Maximum periclase solid solution was found to correspond to 9.5 wt % Al ions and the maximum spinel solid solution to 6 wt % Mg ions [⁴⁵].

The spinel divides the phase diagram into two eutectic systems MgO-MgAl₂O₄ and MgAl₂O₄-Al₂O₄ (Fig. 1). Adjustment of sintering temperature and chemical composition can change the material from a solid solution to a two-phase structure, which improves strength, hardness and crack resistance [⁴⁶]. However, low-purity magnesium aluminate spinel contains considerable amount of impurities like silica and soda. These impurities can form low-temperature eutectics, facilitate liquid-phase sintering and make the spinel structure less homogeneous, with considerable amount of clusters, voids and agglomerations.

In general there are two types of spinel: normal and inverted (reverse). In normal spinels all *A* ions are located in tetrahedral sites and all *B* ions in octahedral coordination. When the structure is inverted, the divalent *A* ions and half of the trivalent *B* ions are located in the octahedral sites, while the remaining *B* ions have tetrahedral coordination. Both normal and inverted spinels have the same cubic Fd3m- structure [⁴⁷]. For an

excellent review of the normal and inverted spinel structures and their mechanical properties see Shukla et al [⁴⁸].

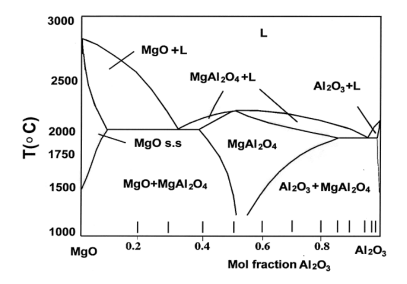


Fig. 1: $MgO-Al_2O_3$ binary phase diagram [⁴⁹].

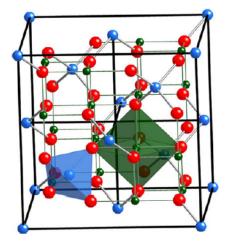


Fig. 2: Tetrahedral cubic $MgAl_2O_4$ spinel unit cell with Mg (blue), Al (green) and O (red) atoms. The unit cell is divided into eight subunits with Mg in tetrahedral (blue) and Al in octahedral (green) coordination in alternating sub-cells [⁴⁸].

2.4 Basic spinel processing

Spinel components are usually fabricated starting from phase-pure spinel powder. In order to enhance sinterability, the powder must be very fine (less than 0.5 μ m in diameter) and have high specific surface area, typically higher than 15 m²/g. Such very fine powders are difficult to process. They tend to have very low bulk density and are difficult to disperse in a liquid medium to facilitate forming processes [⁵⁰].

The first step in spinel powder processing is to spray the powder dry into a denser, more flowable powder containing organic processing aids. Very careful selection of organic additives and mixing methodology are required to successfully disperse a high quantity of powder while retaining sufficient fluidity of the resulting slurry.

Following the spray drying, the powder is molded and pressed, typically using isostatic pressure. This fabrication method is suitable for small to moderate numbers of components ranging in size from small coupons to large plates, and geometries that can include tubes or even domes. Careful control of the process is required to provide sufficient densification in order to minimize shrinkage stresses occurring in later thermal processing [⁵¹].

Pressed parts must be prefired to remove the organic additives prior to sintering. When done incorrectly, this process can lead to the fracture of the pressed part. Consequently, this step of the process requires the longest cycle time of any spinel fabrication process steps.

The sintering process step follows debinding, and the part is densified up to at least 95%. Despite removal of organics in the prior processing step, a relatively long cycle is still required for sintering in order to remove any residual organics, as well as any adsorbed water bound to the very high surface area particles [⁵²].

Hot isostatic pressing (HIP) is used to pressurize the parts during a thermal cycle at temperatures similar to the sintering temperature. Under these conditions, the residual porosity is forced to shrink and ultimately removed as the material densifies further [⁵⁰].

2.5 Physical, chemical and mechanical properties

The basic properties of spinel are governed by the types of atoms present, their bonds and their order. Magnesium aluminate spinel material is based on ionic bonds, and at room temperature fracture occurs without any plastic deformation, which is reflected in a limited tensile strength and toughness [⁵³]. Pores and other imperfections can lead to stress concentrations, further decreasing strength and toughness. Due to the ionic bounding, dislocation slip generally does not occur at room temperature [⁵⁴].

Spinel's optical properties are comparable to polycrystalline alumina and, since it has a cubic structure (i.e. no birefringence), it can transmit light with wavelengths from 0.25 to 5.5 μ m without optical distortion [⁵⁵]. Light transmission through polycrystalline transparent ceramics is influenced by absorption, scattering by pores and additionally decreased by losses due to birefringent splitting of the beam at grain boundaries (Fig. 3). Transmission of light is attained by low porosities (< 0.05 %) along with rather small grain sizes (< 1 μ m) [⁵⁶]. In fact, only 0.1 % residual porosity can completely impair transparency, and large grains increase light scattering [⁵⁷]. MgAl₂O₄ spinel is one of the cubic oxides where high transmission values can be obtained even at larger grain sizes due to the absence of birefringent scattering [²³].

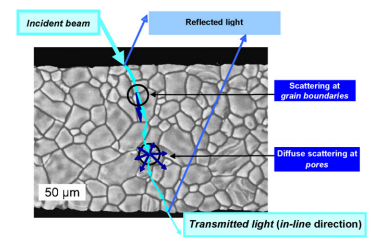


Fig. 3: Light transmission through a polycrystalline translucent ceramic $[^{23}]$.

Mechanical properties of transparent polycrystalline spinels have been reported at room temperature (hardness: 14 - 17 GPa - where applied loads are not always specified, Young's Modulus: 260 - 280 GPa, flexural strength: 185 - 300 MPa and fracture toughness: 1.5 - 2.2 MPa·m^{0.5}) [^{1,46,58,59,60,61}]. Limited data exist for elevated temperature [^{62,63,64,65,66}]. Fracture toughness and Young's modulus of magnesium aluminate spinel as a function of temperature are depicted in Fig. 4. The reduction of elastic modulus and fracture toughness with increasing temperature is typical for many sintered ceramics, especially in cases where the behavior may be governed by a second-phase sintering aid [^{63,67}]. Since for this group of materials such behavior typically coincides with softening of the grain boundaries, a monotonous decrease of mechanical properties with temperature can be expected.

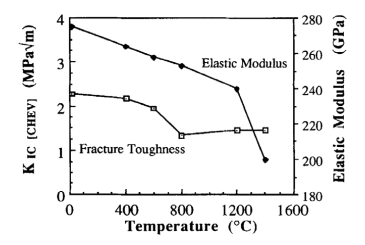


Fig. 4: Fracture toughness and Young's modulus of $MgAl_2O_4$ spinel as a function of temperature $[^{67}]$.

Contrary to Young's modulus and fracture toughness, the hardness of spinel (see section 4.4) is usually load-dependent. However, typically data are only reported for one particular (rather high) load value [⁶⁸]. When a very low indentation load is used, the measured hardness is usually higher. With an increase in test load, the measured hardness decreases (Fig. 5); a behavior often observed for ceramic material, which is known as

indentation size effect (ISE) [⁶⁹,⁷⁰,⁷¹,⁷²,⁷³]. The ISE makes it difficult to define hardness by a single number. Many research works have been devoted to the origin of the ISE and several possible explanations have been proposed. In addition to explanations that concern experimental errors [⁷⁴], elastic recovery work, hardening during indentation, dislocation and twin activities and (especially for ceramic materials), the onset of local damage have been considered [⁷⁵].

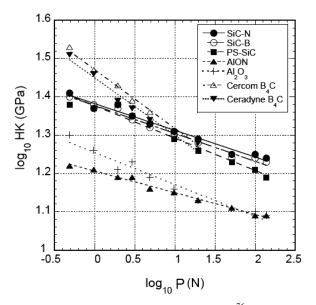


Fig. 5: Knoop hardness-load data for some ceramic materials $[^{76}]$.

Many works have been devoted to improving the mechanical properties of transparent MgAl₂O₄ spinels [51 , 77 , 78 , 79]. Parameters such as non-stoichiometry coefficient *n* and powder composition, as well as the sintering and hot-isostatic pressing (HIP) parameters, significantly changed the mechanical behavior of sintered ceramics [46 , 80]. As illustrated in Fig. 6, the grain size increased from 290 nm (1150 °C) to 9.8 µm (1550 °C) with increasing sintering temperature, while at the same time the porosity slightly decreased.

A variation of the HIP conditions (Table 1) revealed that Young's modulus and fracture toughness are rather insensitive to the grain size. However, increasing HIP temperature and pressure resulted in a decrease of the optical extinction $[^{81}]$.

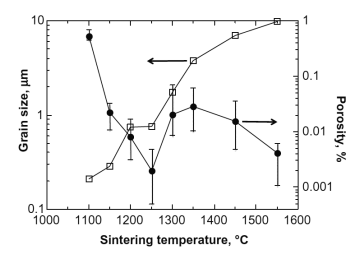


Fig. 6: Dependence of grain size and porosity of $MgAl_2O_4$ spinel on the sintering temperature $[^{80}]$.

Table 1. Basic mechanical properties of $MgAl_2O_4$ spinel as a function of grain size obtained using different processing conditions $[^{82}]$.

Material	Average grain size	Young's Modulus (GPa)	Hardness HV 10 (GPa)	K _{IC} (MPa·√m)	4-point bending strength (MPa)	Compressive strength static – dynamic (MPa)
Spinel ceramics (all transparent)	0.35 μm 0.55 μm 1.4 μm 13 μm (5-50μm) 36 μm (10-75 μm)	$\begin{array}{c} 272 \pm 0 \\ 270 \pm 2 \\ 270 \pm 3 \\ 275 \pm 1 \\ 273 \pm 1 \end{array}$	$\begin{array}{c} 14.7 \pm 0.2 \\ 14.3 \pm 0.2 \\ 13.6 \pm 0.1 \\ 12.7 \pm 0.2 \\ 12.0 - 13.0 \end{array}$	$\begin{array}{c} 1.7 \pm 0.1 \\ 1.7 \pm 0.1 \\ 2.0 \pm 0.1 \\ 1.8 \pm 0.1 \\ 1.8 \pm 0.1 \end{array}$	$\begin{array}{c} 274 \pm 40 \\ 284 \pm 18 \\ 270 \pm 57 \\ 210 \pm 13 \\ 153 - 193 \end{array}$	883 - 3497 879 - 3569 757 - 3510 643 - 3550

Most mechanical properties of non-stoichiometric spinel appear to be superior compared to stoichiometric spinel. However, the flexural strength first increases at lower n and then decreases at higher n, whereas hardness and fracture toughness increase uniformly with increasing n (Fig. 7) [⁴⁶].

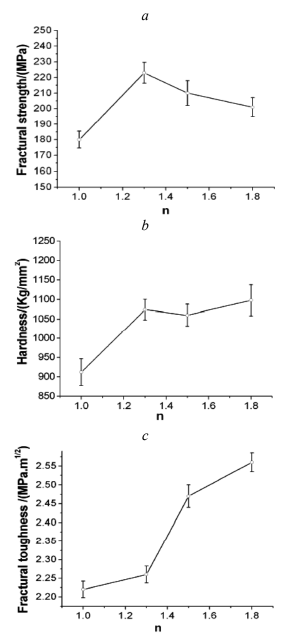


Fig. 7: Mechanical properties of transparent spinel ceramics prepared from MgO- $n-Al_2O_3$ powders as a function of non-stoichiometry n, (a) flexural strength, (b) hardness and (c) fracture toughness [⁴⁶].

The increase of strength might be attributed to better densification related to the enhanced sintering capability of non-stoichiometric ceramics, due to spinel lattice defects induced by deviation from stoichiometry of MgO \cdot nAl₂O₄ powders grains and by corundum grains segregating from spinel grains during cooling. The corundum grains usually improve in strength due to their high elastic modulus and stronger bonding with spinel grains in comparison with the stoichiometric spinel grains [⁸³,⁸⁴]. Hardness increases with increasing *n* due to Al₂O₃ grain separation and the densification of spinel grains [⁸⁵]. Toughening effects are supposed to be increased by dispersed rigid Al₂O₃ particles and non-stoichiometric spinel grains. Rigid Al₂O₃ particles are furthermore able to resist crack expansion and consume energy, thereby improving toughness [⁴⁶].

Furthermore, a clear effect of the porosity on the mechanical properties $[^{39}]$ has been verified (Table 2).

Table 2. Effect of porosity on the mechanical proper	ties of spinel
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Property	Units	Dense MMA (6% porosity)	Porous (18.3%)	Porous (24.5%)
Young's modulus, <i>E</i> Poisson's ratio, v	GPa	192 ± 8	100 ± 10 0.21	70 ± 7
Fracture toughness, K_{Ic} Fracture energy, G_c	$MPa\cdot m^{1/2}$	2.1 ± 0.2 22	1.2 ± 0.09 14	0.63 ± 0.08 5.3
Mean flexural strength, $\sigma_{\rm f}$ Weibull's stress, σ_0 Weibull modulus, <i>m</i>	MPa MPa	151 ± 14 157 10.8	51 ± 6 54 10.7	30 ± 3 31.5 9.3
$c \cong \left(\frac{K_{\mathrm{Ic}}}{\sigma_{\mathrm{f}}}\right)^2$	m	$1.9 imes 10^{-4}$	$5.5 imes 10^{-4}$	$4.4 imes 10^{-4}$
Vickers hardness	GPa	6.3 ± 0.3		

 $(MMA is MgO + MgAl_2O_4)$

In comparison with other industrially used transparent ceramics, spinel can be ranked between sapphire and fused silica (Table 3) [⁸⁶] with regard to its key mechanical properties.

Property	Units	AION	Fused silica	Sapphire	Spinel
Density	kg/m ³	$3.69 imes 10^3$	2.21×10^3	3.97×10^3	3.59×10^{3}
Areal density (at 1" thickness)	kg/m ²	93.89	55.85	100.97	90.86
Elastic modulus	Pa	$334 imes 10^9$	$70 imes 10^9$	$344 imes 10^9$	$260 imes 10^9$
Mean flexure strength	Pa	$380 imes 10^6$	$48 imes 10^6$	$742 imes 10^6$	184×10^{6}
Fracture toughness	Pa m ^{1/2}	$2.4 imes 10^6$	$0.78 imes 10^{6}$	$3.0 imes 10^6$	$1.7 imes 10^6$
Knoop hardness	Pa	$17.7 imes 10^9$	$4.5 imes 10^9$	19.6×10^9	14.9×10^{9}
Transmission in visual spectrum	%	82–85	91–92	75–82	I
Maturity of technology		Relatively new technology (becoming commercially viable)	Well established technology	Well established technology	Established, continued advancements
Cost		3-5 times that of glass	Lowest material and processing costs	Higher than ALON TM	Lower than ALON TM
Manufacturing costs		High due to high processing temperature, proprietary powder, and surface finish	Relatively low due to lower melting temps	High due to high temperature processing and surface finish requirements	Moderate due to surface finish requirements
Bottleneck		requirements Cost and limited dimensions	Limited hallistic	Cost and limited	Limited dimensions
			protection enhancement	dimensions	
Commercial availability		Sumert Corp., limited availability	Widely available	Widely available in smaller sizes	In the process of becoming more commercially available
Environmental resistance		Low chemical reactivity and highly scratch resistant		Low chemical reactivity and highly scratch resistant	

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Table 3. Typical material properties of transparent ceramics [⁸⁶].

LITERATURE REVIEW

2.6 Mechanical characterization

2.6.1 Young's Modulus

2.6.1.1 Indentation

In contrast to traditional hardness testing, instrumented depth sensitive indentation permits the measurement of the penetration depth of the indentation tip into the material h as a function of force P (Fig. 9). For indentation testing a Vickers tip is most commonly used. The Vickers indenter geometry corresponds to a square pyramid, with an angle of 136° between the triangle faces. Indentation load-displacement curves can be used to determine mechanical characteristics, according to DIN 50359-1 [⁸⁷] (Fig. 8).

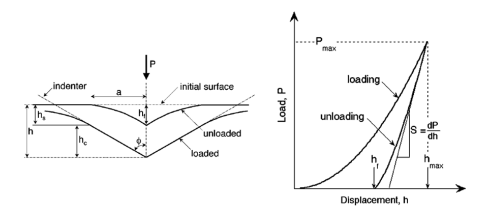


Fig. 8: Instrumented indentation: a) main parameters describing the impression, b) schematic representation of load - displacement curve obtained during indentation $[^{88}]$.

The reduced Young's modulus can then be calculated from the indentation loaddisplacement curve [⁸⁸]:

$$E_r = \beta \frac{\sqrt{\pi}}{2} \cdot \frac{S}{\sqrt{A_c}}$$
 Eq. 1

where β is a correction factor for the Vickers indenter (1.0124); *S* is the unloading slope dP/dh and A_c is the projected contact area predetermined via calibration of the indentation system. The indentation Young's modulus can then be calculated as [⁸⁸]:

$$E_{IT} = \frac{1 - v^{2}}{\frac{1}{E_{r}} - \frac{1 - v_{i}}{E_{i}}}$$
 Eq. 2

where E_i is Young's modulus of the indenter, and v and v_i are Poisson's ratios of the tested material and indenter, respectively.

2.6.1.2 Bending test

In the case of a bi-axial ring-on-ring bending test, the Young's modulus is calculated from the linear part of the load-displacement curve after ASTM C1499-05 [⁸⁹]:

$$E_{ROR} = \frac{3 \cdot \left(1 - \nu^2\right) \cdot r_1^2 \cdot \Delta P}{2\pi \cdot \Delta f \cdot h^3} \times \left[\left(\frac{r_2}{r_1}\right)^2 - 1 - \ln\left(\frac{r_2}{r_1}\right) + \frac{1}{2} \left(\frac{1 - \nu}{1 + \nu}\right) \times \left(\frac{r_2^2 - r_1^2}{r_3^2}\right) \times \left(\frac{r_2^2}{r_1^2}\right) \right] \text{ Eq. 3}$$

where v is Poisson's ratio, ΔP is the difference in force, and Δf is the corresponding change in displacement, h is the specimen thickness, and r_1 , r_2 and r_3 are radii of load ring, support ring and specimen, respectively.

2.6.1.3 Impulse excitation technique

The impulse excitation technique (IET) is a non-destructive method to determine the Young's modulus, with widespread use in science and industry. This method is based on an actuator that hits the surface of the sample, resulting in a free vibration with a material-specific and geometry-dependent resonance frequency. Since every sample has its own resonance frequency and the geometry is fixed, the resonance frequency is only related to density and the Young's modulus [90]. Often the entire oscillation waveform after excitation is recorded, as illustrated in Fig. 16. Afterwards, the fundamental frequency f of the flexural resonance is extracted automatically.

The Young's modulus is often calculated from resonance frequency, using automatized software based on the relevant relationships for disc- and bar-shaped specimens given in ASTM E1876 - 01.

This method's advantage lies not only in the simplicity of the experimental setup, but also in high measurement accuracy. In fact, it has been verified for glass that the Young's modulus measured by bending tests is usually ~7 % lower than that obtained using IET, due to system error and contact displacement [⁹¹]. IET results have excellent stability, while results of Vickers indentation show bigger scatter and are influenced by sample surface and inhomogeniety effects [⁹¹].

2.6.2 Hardness

Additionally, the depth-sensitive indentation technique permits the determination of indentation hardness (H_{IT}) by the ratio of maximum load to residual contact area [⁸⁷]:

$$H_{IT} = \frac{P_{\text{max}}}{A_C}$$
 Eq. 4

whereas the Vickers hardness can be determined as:

$$HV = \frac{2P\sin\frac{136^{\circ}}{2}}{d^{2}} \approx 1,854\frac{P}{d^{2}}$$
 Eq. 5

where *d* is the arithmetic mean of the two diagonals.

2.6.3 Fracture toughness

Fracture toughness (K_{IC}) is the material resistance to instable crack growth. Fracture toughness can be tested by different methods, e.g., single edge notch bending, indentation and indentation-strength-in-bending tests [⁴⁰], and by means of the Griffith criterion. The following sections give details on these methods.

2.6.3.1 Indentation

The local contact of a sharp tip with the surface of brittle materials can result in damage by crack formation at the edges of the indenter tip. The size of the impression and cracks is related to the elastic/plastic stress field under the indenter. If a material shows indentation cracking, the fracture toughness can be determined, defined here as the indentation toughness (K_{IND}). Different relationships exist to determine indentation toughness, depending on the crack types generated during the impression. Evans and Charles [⁹²], Niihara et al. [⁹³], Anstis et al. [⁹⁴] and Lawn et al. [⁹⁵] have been pioneers in determining fracture toughness by Vickers indentation. The fracture toughness is computed from the length of the indentation-induced cracks, taking into account applied load, hardness, Young's modulus and calibration geometry factor [^{94,95,96}]. Due to the small size of indents and cracks, this method is a macroscopically rather nondestructive characterization. However, the obtained values need careful interpretation. Mistakes in crack length observation and irregular fracture behavior can result in errors of up to 50 % [⁹⁷]. The typical crack pattern after Vickers indentation is given in Fig. 9.

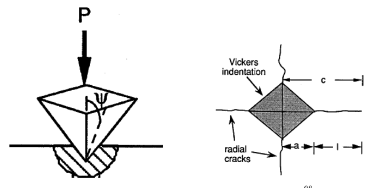


Fig. 9: Radial crack system (right) after Vickers indentation (left) [98].

In the present study, the widely used relationships for a half-penny crack system proposed by Lankford [⁹⁹], Niihara [⁹³] and Anstis [⁹⁴] were applied for indentation toughness determination.

Lankford equation [⁹⁹]:
$$K_{ind} = 0.0363 \cdot (E/H)^{2/5} \cdot (P/a^{1.5}) \cdot (a/c)^{1.56}$$
 Eq. 6
(c/a > 2)

Niihara equation [⁹³]:
$$K_{ind} = 0.067 H \sqrt{a \cdot (E/H)^{0.4} \cdot (c/a)^{-1.5}}$$
 Eq. 7
(c/a > 2.5)

Anstis equation [⁹⁴]: $K_{ind} = 0.016 \cdot (E/H)^{0.5} \cdot (P/c^{1.5})$ Eq. 8 ($c \ge 2a$)

where: a – half of imprint's diagonal length, c – crack length (measured from the end of imprint) and P – applied load.

2.6.3.2 Indentation-Strength-Method

An additional method of determining fracture toughness is the Indentation-Strength-in-Bending method (ISM). This method is based on the fracture of pre-indented specimens. Consequently, well-defined defects are used to determine the fracture toughness from the fracture load obtained in a bending test. The fracture toughness (K_{ISM}) can be calculated using [⁹⁶]:

$$K_{ISB} = A \cdot \left(\frac{E}{H}\right)^{\frac{1}{8}} \cdot \left(\sigma \cdot P^{\frac{1}{3}}\right)^{\frac{3}{4}}$$
 Eq. 9

During the bending experiment, the indentation induced residual stress field results in subcritical (stable) growth of the indentation-induced crack. This effect is considered by the geometrical constant $A = 0.59 \pm 0.12$ [⁹⁴,⁹⁶], which is determined as $A = \left[(256/27) \cdot (\pi \cdot \Omega)^{\frac{3}{2}} \cdot \varphi \right]^{\frac{1}{4}}$, where Ω is a crack-geometry factor considering free-surface effects (ellipticity and crack interaction) [⁹⁶,¹⁰⁰], and $\varphi = 0.016 \pm 0.004$ is a

material independent "calibration" constant for radial cracks at Vickers indents, which was obtained by averaging over the data for structured ceramics [⁹⁴,⁹⁵]. This calibration value is calculated from the ratio between the maximal crack length after stable crack growth c_m (just before catastrophic specimen failure) and the initial indentation-induced crack length c_0 which was $\Omega \sim (c_m/c_0)^{3/8} = 1.12$. The calibration value needs to be verified experimentally for any material and if necessary *A* needs to be re-calibrated.

2.6.3.3 Single edge notch bending

Fracture toughness of ceramics is often determined by SENB tests, according to ISO 13586 [¹⁰¹]. This method, based on a notched bending specimen, is one of the simplest and most reliable methods for determining fracture toughness [⁴⁰].

As illustrated in Fig. 10, an edge crack of depth *c* that is pre-introduced by cutting and sharpened with a razor blade, is extended from the tensile surface when a critical load is reached. In case of four-point bending of rectangular bar-shaped specimens, the critical stress intensity factor (fracture toughness) is then given by $[^{102}]$:

$$K_{SENB} = \frac{P_{C}}{B\sqrt{W}} \cdot \frac{S_{1} - S_{2}}{W} \cdot \frac{3\Gamma_{M}\sqrt{\alpha}}{2(1-\alpha)^{3/2}}$$
 Eq. 10

where P_c is the critical fracture load, *B* is a thickness, *W* is a height, S_2 inner and S_1 outer roller span, and Γ_M is numerical factor:

$$\Gamma_{M} = 1.1215\sqrt{\pi} \cdot \left[\frac{5}{8} - \frac{5}{12}\alpha + \frac{1}{8}\alpha^{2} + 5\alpha^{2}\beta^{6} + \frac{3}{8}\exp\left(-6.1342\frac{\alpha}{\beta}\right)\right]$$
 Eq. 11

where $\alpha = a/W \approx 0.5$ and $\beta = 1 - \alpha$.

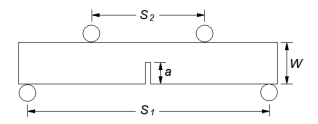


Fig. 10: Four-point bending test with edge-notched specimen \int_{0}^{40}].

2.6.3.4 Griffith criterion

In bending tests, failure will occur from pre-existing flaws. Therefore, fracture toughness determination is possible if these flaws are identified and their size is measured. If enough fracture energy is available, which is usually the case for finite-sized specimens in a bending test, fracture occurs when a critical applied tensile stress is exceeded. In this case the stress intensity factor exceeds the fracture toughness and unstable crack growth starts [$^{103}, ^{104}, ^{105}$]. The major types of fracture-initiating defects in ceramics are machining flaws, microstructural inhomogeneities, inclusions and pores. Fracture toughness is related to the critical stress and defect size *c* via:

$$K_{IC.Griffith} = (Y/Z) \cdot \sigma \cdot \sqrt{c}$$
 Eq. 12

where *Y* is a geometry factor dependent on the loading conditions, specimen shape and defect size, and *Z* is a shape factor for the defect. For flaws in a uniform stress field, *Y* is independent of *c* and given by: $Y = \sqrt{\pi}$ - for volume flaws, and $Y = 1.12 \cdot \sqrt{\pi}$ - for surface flaws. If the defect has a linear crack front, the shape factor is Z = 1, whereas for a circular or penny-shaped flaw, $Z = \frac{\pi}{2}$ [⁹⁸].

Therefore, an estimation of fracture toughness is possible based on experimental observations of fracture origins using fractography.

2.6.4 Strength

Fracture stress is determined in the case of ring-on-ring bending tests from the maximum load at failure *P* (see section 3.3.2), according to ASTM C1499-05 [⁸⁹]:

$$\sigma = \frac{3(1+\nu)P}{2\pi \cdot h^2} \times \left[\ln \frac{r_2}{r_1} + \frac{(1-\nu)}{(1+\nu)} \times \frac{r_2^2 - r_1^2}{2 \cdot r_3^2} \right]$$
Eq. 13

where v is Poisson's ratio, h is the specimen thickness, and r_1 , r_2 and r_3 are radius of load ring, support ring, and specimen radius, respectively.

Fracture stress of ceramics typically displays a considerable scatter that depends on the defect size distribution. Strength characterization is therefore usually performed using statistical approaches. In particular, Weibull statistics are widely used to assess the inherent scatter [¹⁰⁶,¹⁰⁷]. Based on the "weakest link hypothesis", it is assumed that the most serious (largest) flaw controls the strength [¹⁰⁸,¹⁰⁹]. In general, the critical parameters for predicting the fracture strength of a component are the specific dimensions, characteristic strength, Weibull modulus and threshold strength.

According to the Weibull statistics that are widely accepted for brittle materials [¹⁰⁸], the cumulative failure probability of a component made of brittle material with specified dimension $P(\sigma)$, subjected to stress σ , is given by:

$$P(\sigma) = 1 - \exp\left[-\left(\frac{\sigma - \sigma_u}{\sigma_0}\right)^m\right]$$
 Eq. 14

where σ_0 is the characteristic fracture strength, σ_u is the threshold stress, below which no failure will occur, and *m* is the Weibull modulus, being a measure of strength distribution.

The characteristic strength corresponds to a failure probability of 63.2%, and therefore is only a weak criterion for assessing the reliability of brittle materials. Only the knowledge of characteristic strength, Weibull modulus and threshold strength permits complete characterization of a material for a given specimen dimension and an estimation

of the probability of failure for a particular stress level. Often σ_u is assumed to be zero, yielding a two-parameter relationship. In fact, numerical simulations and experimental data have verified that as long as sample data are limited in number (~ 30) and the threshold stress is not too large, a two-parameter Weibull distribution should be used [¹¹⁰,¹¹¹]. It has also been demonstrated that when a two-parameter distribution is used, an underestimation is more likely than an overestimation. Hence, it can be considered the more conservative approach [¹¹²]. The statistical parameters of the Weibull distribution are most frequently assessed using linear regression (LR) or the maximum likelihood method (ML) [¹¹³]. In the present study, the LR method is used for simplicity, since it has been verified that the differences in the resulting parameters obtained using the different methods is rather small [¹¹⁴]. Obviously the Weibull parameters are subject to uncertainties and, for design purposes, it is necessary to give appropriate confidence intervals for example based on ASTM C1239 – 06A [¹¹⁵].

In the linear regression analysis (LR) the least-square fitting of a two-parameter Weibull distribution, which takes twice the double logarithm of Eq. 14, permits to determine *m* as the slope, and σ_{θ} as the y-intercept:

$$\ln \ln \left(\frac{1}{1 - P_f} \right) = m \cdot \left(\ln(\sigma) - \ln(\sigma_0) \right)$$
 Eq. 15

where P_f is the failure probability. In the LR method, the stress values are ranked with respect to their individual probability according to [¹¹⁶]:

$$P_i(\sigma) = (i - 0.5)/N$$
 Eq. 16

Since the characteristic strength is a rather inappropriate parameter to assess the reliability of a brittle component (given the failure probability of 63.2%), usually fracture stresses for failure probabilities of 10^{-6} or 10^{-3} should not be exceeded to warrant structural integrity. Furthermore, the decrease of fracture stress due to larger deformed volumes in real application should be considered.

In addition to the two-parameter distribution, a three-parameter Weibull distribution can also be used to analyze the data $[^{117}]$. In order to carry out the linear

regression analysis, a threshold stress is chosen and the parameters are optimized based on the uncertainty of the regression slope.

The standard deviation of Weibull modulus *s* (*m*) and characteristic strength *s* (σ_0) can be estimated for a particular number of tests *g* via [¹¹⁸]:

$$\frac{s(m)}{m} = \frac{1}{\sqrt{g}}$$
 Eq. 17

$$\frac{s(\sigma_0)}{\sigma_0} = \frac{1}{m \cdot \sqrt{g}}$$
 Eq. 18

The statistical bias errors in the estimated Weibull parameters and calculation of confidence bounds have to be calculated from the available data set [¹¹⁹]. Depending on the number of specimens in a given batch, the estimate of Weibull modulus *m* may exhibit significant statistical bias. An unbiased estimate of the Weibull modulus m_U is obtained by multiplying the biased estimate by an appropriate unbiasing factor (listed in ASTM C1239-06A). For this procedure, an unbiasing factor of $1-1.593145 N^{-1.046958}$ suggested in [¹¹⁶] can also be applied, yielding a negligible difference of ~ 2%, compared with the tabulated ASTM values for a data set of at least 10 specimens.

The equations for the upper $m_{UPP} = m_{BLAS} / m_5 \%$ and lower $m_{LOW} = m_{BLAS} / m_{95} \%$ confidence intervals, respectively, describe how the different limiting values are calculated for a 90% confidence level. The values for $m_5 \%$ and $m_{95} \%$ are listed in ASTM C1239-06A [¹¹⁵,¹²⁰].

The strength values obtained using ring-on-ring specimens vary with test specimen size. Characteristic strength values can be scaled to other specimen geometries using the area relationship (Eq. 19). The effective area under tensile stress in a ring-on-ring test is [¹²¹,¹²²]

$$A_{eff.ROR} = \frac{\pi}{2} D_{L}^{2} \left\{ 1 + \left[\frac{44(1+\nu)}{3(1+m)} \right] \cdot \left[\frac{5+m}{2+m} \right] \cdot \left(\frac{D_{s} - D_{L}}{D_{s}D} \right)^{2} \left[\frac{2D^{2}(1+\nu) + (D_{s} - D_{L})^{2}(1-\nu)}{(3+\nu) \cdot (1+3\nu)} \right] \right\} \text{ Eq. 19}$$

Comparison of Eq. 19 to numerical computations indicates that for $m \ge 5$, $v \ge 0.17$ and $D_s / D_L = 0.5$, the error is less than 3% compared to a full integration of the stress field, but for $D_s / D_L = 0.2$, the estimates are better than 5%. The use of Eq. 19 requires that fracture origins are clearly defined as surface defects [¹²¹].

Since the strength of ceramics depends on the surface area or volume under stress, the characteristic strength needs to be rescaled from the size (effective area) of the test specimen to the actual component via [123]:

$$\left(\frac{\sigma_{0,G1} - \sigma_u}{\sigma_{0,G2} - \sigma_u}\right) = \left(\frac{A_1}{A_2}\right)^{-1/m}$$
 Eq. 20

Similar relationships can be derived for volume defects.

2.6.4.1 Local strength

Hertzian (spherical) indentation of brittle materials is a feasible way to test the contact damage that may occur in application. An advantage of Hertzian indentation is that the material first responds linear-elastically before permanent contact damage (fracture) starts. Ring cracking is caused by the radial tensile stress occurring on the edge of the circular contact area, between the indenter and investigated material (Fig. 11).

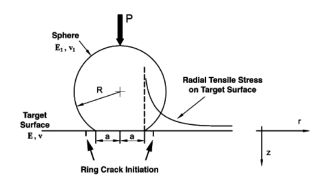


Fig. 11: Schematic diagram for Hertzian indentation and ring crack initiation [124].

When a hard sphere (radius *R* and elastic constants E_1 , v_1) is pressed with a load *P* on to a flat substrate (elastic constants E_2 , v_2), the contact radius *a* is given by $[^{125}]$:

$$a = \sqrt[3]{4kP \cdot r/3E}$$
 Eq. 21

where

$$k = (9/16) \cdot \left[(1 - v_1^2) + (1 - v_2^2) \cdot E_1 / E_2 \right]$$
 Eq. 22

Considering the average stress criterion as a good indicator of material strength, the following relation has been proposed for calculating the local fracture strength [¹²⁵]:

$$\sigma_{Hertzian} = \left[-0.1373a_c^2 + 0.2862a_c + 0.0236 \right] \cdot \left(\frac{3E}{4k}\right)^{\frac{2}{3}} \cdot \frac{1}{\pi} \cdot \left(\frac{P_c}{r}\right)^{\frac{1}{3}}$$
 Eq. 23

where a_c can be determined by Eq. 21 with $P = P_c$.

The effective area ($A_{eff:Hertzian}$) is located at the specimen surface, where z = 0 and corresponds to the region where tensile surface stresses occur. This region is located outside the contact area. $A_{eff:Hertzian}$ is derived for the conditions: $r \ge a$ and z = 0 [¹²⁴] (see Fig. 11).

$$A_{eff.Hertzian} = \frac{\pi}{m-1} \cdot \left(\frac{4kPR}{3E}\right)^{\frac{2}{3}}$$
 Eq. 24

2.6.5 Statistical lifetime evaluation

2.6.5.1 Slow crack growth and Strength-Probability-Time (SPT) prediction

A ceramic component must be designed not only with respect to the applied load, but also to sustain a certain lifetime. For this reason, knowledge of the relationship between strength and operation time is necessary. Existing cracks may grow slowly, depending on the atmosphere, especially humidity. This behavior is known as subcritical crack growth (SCG). If SCG is described quantitatively, the lifetime of the component can be predicted using a strength-probability-time (SPT) diagram. Accordingly, an SPT diagram is a link between strength, probability of failure, and lifetime.

The experimental determination of an SPT diagram can be based on measurement of strength as a function of the loading rate, since a slower loading rate permits cracks to grow during loading, resulting in lower apparent strength. The fracture strength is in then correlated with the stress rate $\dot{\sigma}$ via [¹¹³]:

$$\log \sigma_0 = \frac{1}{n+1} \log \dot{\sigma} + \log D$$
 Eq. 25

where n and D (in MPa/s) are the SCG parameters.

To apply the SPT method, fracture stresses for a monotonous loading need to be known as a function of loading rate and these stresses are condensed to the equivalent stress σ_{1s} . The characteristic strength σ_0 , measured at a certain stress rate of $\dot{\sigma}_1$ is in this case converted into an equivalent stress that would have caused the sample to fail in a time of 1s with a fracture probability of $P_f = 63.2 \% [^{126}, ^{127}]$:

$$\sigma_{1s} = \sigma_0 \left(\frac{\sigma_0}{\dot{\sigma}(n+1)} \right)^{\frac{1}{n}}$$
 Eq. 26

The combination of this stress and the failure probability of 63.2% results in a first data point on the SPT diagram. The 1s line crosses the σ_{Is} data point and has a slope equal to the Weibull modulus *m*. The prediction of different lifetimes can be performed using the relation:

$$\frac{t_2}{t_1} = \left(\frac{\sigma_1}{\sigma_2}\right)^n$$
 Eq. 27

Therefore, a series of lines can be drawn, with spacing between the lines equal to (ln10)/n. In this case, each line represents a one-decade increase in lifetime. Eq. 2 can

then be used to assess failure probability as a function of stress for a given geometry. To analyze the influence of the specimen or component size on the allowable stress, Eq. 20 should be used.

The SPT diagram consists of a series of failure lines representing equivalent times of failure at different stresses. An example for yttria-stabilized zirconia (YSZ) is given in Fig. 12 [¹²⁸].

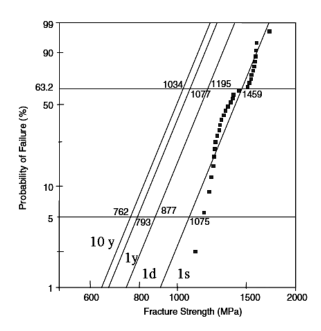


Fig. 12: Strength-probability-time diagram for YSZ [¹²⁸]

It can be seen that for a given failure probability, YSZ survives for a longer period at lower stresses. For example, for a lifetime of 1s with a survival probability of 63.2%, the stress should not exceed ~ 1460 MPa, whereas for a lifetime of 10 years with the same survival probability, the applied stress should not exceed ~ 1030 MPa.

It has to be emphasized that the SPT diagram is only valid for the actual environmental conditions under which the specimens are tested $[^{129}]$.

2.6.6 Photoelasticity

Photoelasticity is a non-destructive experimental method used primarily for two dimensional plain strain state analysis, which uses optical effects to determine strain distribution. Photoelastic analysis provides qualitative assessment of highly strained areas and peak strain at the surface and in the interior of the structure. It also distinguishes areas of low strain level where the material is less utilized. Almost all transparent isotropic materials (such as glass, polycrystalline ceramics, as well as many transparent synthetic resins and polymers) exhibit a double refracting effect on a beam of light, when they are subjected to strain [¹³⁰].

When the material is strained and a ray of light enters along one of the directions of principal strain, the light splits into two component waves – each with its plane of polarization parallel to one of the remaining two principal planes (planes on which shear stress is zero). Furthermore, the light travels along these two paths with different velocities, which depend on the magnitudes of the remaining two principal strains in the material. The incident light is resolved into components having polarization planes parallel to the directions of the principal strains σ_1 and σ_2 (Fig. 13). Since these waves intersect the body with different velocities, the waves appear with a phase relationship, or relative retardation. Quantitative determination of the strain values is usually not possible and the method is used only qualitatively [¹³¹].

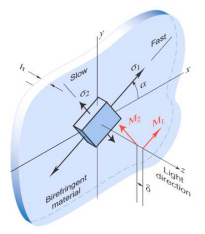


Fig. 13: Birefringent effect [¹³¹]

3 INVESTIGATED MATERIALS AND EXPERIMENTAL METHODS

3.1 Material

3.1.1 Processing

The material was developed, produced and supplied by CeramTec-ETEC, Lohmar. A nano-sized MgAl₂O₄ spinel powder with a specific surface area BET of 30 m²/g, average particle size 70 nm and purity of more than 99.9% was used as raw material. This powder was dispersed with a stirred media mill in water to form a slurry with 50 wt.-% solid material. Polyvinyl alcohol and Ammonium stearate were used as plasticifiers. The slurry was spray dried and then uniaxially pressed on a Herzog HP40 laboratory press. Thereafter, the green samples were debindered, sintered in a Nabertherm HT40 furnace and afterwards hot isostatically pressed in an EPSI Hot Isostatic Pressing. A detailed description of the fine- and coarse-grained spinel production is given in Table 4.

Fine-grained spinel	Coarse-grained spinel
Mixing water, powder (50 wt-%),	Mixing water, powder (50 wt-%),
dispersant, sintering aid	dispersant, sintering aid
Ball milling (2 h)	Ball milling (2 h)
Spray drying (100 – 200 µm)	Freeze drying $(100 - 200 \mu m)$
Molding	Hot pressing (1700 °C, 1 h)
Debindering (24 h / 800 °C)	Hot isostatic pressing (1600 °C, 200 MPa, 2 h)
Sintering (1600 °C / 2h)	
Hot isostatic pressing (1600 °C, 200 MPa, 2 h)	

Table 4. S	Spinel pr	oduction
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3.1.2 Specimens and surface preparation

The transparent spinel was produced as $100 \text{ mm} \times 100 \text{ mm}$ plates, with a thickness of 6.2 mm. Disk shaped specimens were cut out of the spinel plates by wire cutting (Fig. 14) for mechanical testing.

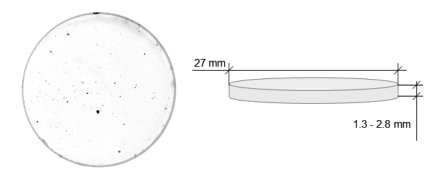


Fig. 14: Disk shaped specimens

Grinding conditions were chosen according to initial results of batches grinded under the different conditions: 54 μ m diamond abrasive wheel (1 min), 18 μ m diamond abrasive wheel (3 min) and final polishing using 6 μ m diamond suspension (30 min). The optimal polishing condition (6 μ m) was selected by CeramTec-ETEC to obtain the required optical properties, whereas the optimal grinding condition (18 μ m) was chosen after macromechanical measurements (see section 4.7). These specimens were investigated by ring-on-ring bending and impulse excitation. The samples were polished (Minimet 1000) using diamond paste (1 μ m) and silica slurry (0.2 – 0.5 μ m) for microscopic analysis and indentation testing.

The fine- and coarse-grained spinel specimens investigated by various methods at room (RT) and high temperature (HT) (see section 3.1.3) are given in Table 5.

Method	Fine-grained spinel (number of specimens)		Coarse-grained spinel (number of specimens)	
Methou	Room temperature	High temperature	Room temperature	High temperature
Fractography and microstructure observation	45	11 (etched)	13	4 (etched)
Photoelasticity	16	2 (annealed)		
In-situ fracture observation in light microscopy	1			
Determination of volume and agglomeration-size distribution	25			
X-ray analysis	1		1	
Ring-on-ring bending test	170		52	
Micro- and macro-indentation testing	17	3 (annealed)	14	2 (annealed)
Impulse excitation testing	3	1	3	1
Indentation (K _{IND}) toughness evaluation	9		3	
Single edge notch bending (K _{SENB}) toughness evaluation	5		3	
Indentation strength in bending (K _{ISM}) toughness evaluation	4			

Table 5. Spinel specimens tested in this work

3.1.3 Thermal treatment

The heat treatments of specimens used for microscopic investigations are summarized in Table 6.

Table 6. Application of the heat treatment regimes.

Investigation	Applied methods	Heat treatment	Samples
Effect of annealing on local strains	Photoelasticity	annealing in air [*] (1000 °C, 60 min)	disk-shaped, fragments
Structural observation	Light and scanning electron microscopy	etching in air (1500 °C, 30-120 min)	fragments
Effect of annealing on mechanical properties	Bending, indentation	annealing in air [*] (1000 °C, 60 min)	disk-shaped, fragments

* heating rate 3 K/min, cooling rate 5 K/min.

Annealing conditions were chosen to retain transparency of fine- and coarsegrained spinel samples.

Thermal etching was carried out in order to examine the grain structure. Since sintering was carried out at a temperature of 1600 °C, a temperature 1500 °C was chosen for thermal etching as suggested by the manufacturer. The spinel structures showed different grain size-dependent etching rates. Therefore, exposure times of 30 and 120 min were used for the fine- and coarse-grained spinels, respectively.

3.2 Microstructural observation

3.2.1 X-ray diffractometry

The phase composition was determined by X-ray diffraction using a Siemens D5000 diffractometer, equipped with an X-ray tube with copper anode ($\lambda = 1.5418$ Å), powered with an accelerating voltage of 40 kV and a current of 40 mA.

3.2.2 Microscopy

Microstructure investigations were performed by light microscopy (LM: Zeiss Axiomat), confocal laser scanning microscope (CM: Keyence VK-9500) and scanning electron microscopy (SEM: LEO 440) on polished, ceramographically prepared cross-sections. Specimens were embedded in epoxy resin and ground on silicon carbide abrasive papers with stepwise decreasing size of abrasive particles (400 up to 4000 grit). For subsequent polishing, diamond pastes and suspensions with grain sizes from 3, through 1 to 1/4 µm were used. When visibility of the grain structure was required, the epoxy resin was removed and the polished ceramographic sections were thermally etched (Table 6).

Image analysis (grain size, porosity and phase content estimation) was carried out using the "AnalySIS" software, provided by Olympus., Since thermal etching did not appear to influence the grain structure, grain size analysis was performed on thermally etched specimens (although a local influence on the phase composition could not be ruled out). The latter, however, is not critical for grain size analysis. Grain sizes were determined from light and scanning electron micrographs. The planimetric method was applied involving automated measurement of the grain area for grain size determination, and was performed by the AnalySIS software. The results were then recalculated into the diameter of a circle with an identical area as the measured grain, and are given in the following as an equivalent circular diameter (ECD). The average grain size of the samples was calculated, assuming spherical grains, from the linear intercept on a two-dimensional cross-section using the following relation [¹³²]:

 $D = 1.56 \cdot D_1$

Eq. 28

where D is the corrected grain size and D_I is the average grain size obtained from the linear intercept method.

A light microscope (Leica DMLM) with a magnification of 63 was used at CeramTec-ETEC for analyzing intrinsic agglomerates, as well as providing agglomerate dimensions and distribution. In order to determine sizes and volume of the particles (Fig. 45), an automatical recalculation into white/black images was obtained by digital images taken from the center of the specimen ($\sim 200 \text{ mm}^2$), as well as carried out by single-purpose software.

To reveal fracture mode and crack shape of the fine- and coarse-grained spinels, a line of Vickers indents was introduced into the surfaces of some samples under a load of 49 N, with a distance of $\sim 20 \ \mu m$ between the radial crack tips. Subsequently, the specimen was manually broken and observed via light microscopy.

Fractographic analysis was performed on specimens fractured in bending tests. Fracture origins were identified by a stereo-zoom microscope (Olympus SZH10) and SEM (LEO 440), as well as by a confocal laser-scanning microscope (Keyence VK-9500). The SEM in combination with Energy Dispersive X-ray spectroscopy (EDX) at high resolution (1 μ m) permitted to analyze the elemental composition.

Transmission electron microscopy (TEM: Philips CM 200 and JEOL 200CX) was conducted on thin foils prepared by Ga^+ - focused ion beam technique (FIB: Zeiss Neon 40EsB CrossBeam). Critical microstructural details were investigated by means of high-resolution TEM.

The photoelastic observation was carried out with the aid of a circular polariscope in dark field configuration (Zeiss Axioskop 40 Pol). The set-up consists of a light source, a polarizer, a quarter-wave plate oriented at 45° with respect to the polarizer, the specimen, a second quarter-wave plate, and an analyzer (Fig. 15).

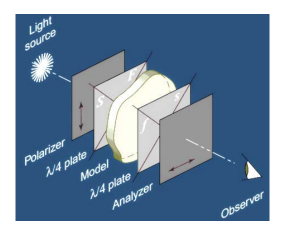


Fig. 15: Working scheme of circular polariscope (dark field) [¹³¹]

3.3 Mechanical characterization

Various testing methods were applied to obtain mechanical properties of the MgAl₂O₄ spinel. In particular, bending tests were used to determine fracture strength and Young's modulus, which was additionally determined using a resonance-based method and depth-sensitive indentation. Fracture strength data were statistically evaluated; fractured specimens were used to assess the failure origin. The depth-sensitive indentation method was also used to measure the hardness of the material. Fracture toughness was evaluated by three methods: the length of indentation cracks, the indentation strength method and from a direct measurement of the defect size in combination with the Griffith criterion.

3.3.1 Impulse-excitation technique

The measurements at room temperature were carried out on disk-shaped specimens with a 27 mm diameter and thicknesses between 1.3 and 2.8 mm (see section 3.1.2), using a commercial resonance system (GrindoSonic, Lemmens KG, Belgium) (Fig. 16a) based on ASTM E1876-01 [¹³³]. However, for high temperature measurement the use of rectangular bars was required by the test setup (25 mm × 6 mm × 2 mm); the bars were wire cut out of disk shaped specimens.

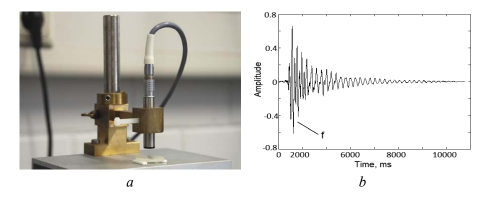


Fig. 16: a) Set-up for impulse excitation tests in air; *b)* typical resonance signal of the specimen induced by push rod.

3.3.2 Bending test

Bending experiments were carried out using the electromechanical testing machine Instron 1362 (Fig. 17a). Specimens for strength characterization were investigated in a ring-on-ring configuration, which avoids edge effects, and failure initiates within the loading ring. A limited number of specimens have been machined to carry out the single edge notch bending (SENB) test after ISO 13586 (see section 3.5.3) to assess the fracture toughness.

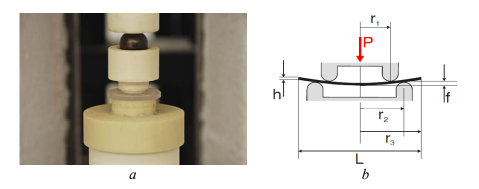


Fig. 17: a) Ring-on-ring testing set-up; b) Schematic sketch of load and support ring with specimen.

Biaxial ring-on-ring bending test, according to the procedure given in ISO 1288 [¹³⁴], permits a simultaneous assessment of Young's modulus and fracture stress for thin plates. The specimens were mounted between a loading (top) and support ring (bottom), both made of alumina. Alumina ceramics were also used for the upper pushing rod and the lower tubular support piston. The arrangement provided high temperature stability, stiffness, and good alignment of the load train. The displacement was measured in the center of the specimen with a ceramic extension rod attached to a linear variable differential transformer (Sangamo, LVDT, range ± 1 mm, precision 1.25 µm). The load was determined with a 10 kN load cell (Interface 1210 ACK). The temperature was monitored during the test close to the outer specimen surface with a thermocouple. All tests were carried out with support and load ring diameters of $D_s = 19$ mm and $D_L = 9.4$ mm, respectively. Young's modulus was measured at specimens with thicknesses ranging from 0.5 mm to 3 mm. In order to determine fracture stress, 3 mm thick specimens were used.

To determine inert strength, a loading rate of 100 N/min was chosen in order to minimize slow crack growth effects. Ten tests per rate at stress rates of $3.6 \cdot 10^2$, $3.6 \cdot 10^1$, and 19 tests at 3.6 MPa/s were performed to analyze slow crack growth.

Limitations for the use of the ring-on-ring test are provided in the literature $[^{89,121}]$. The linear theory is valid as long as the deflection of the specimen does not exceed a certain value, which depends on the diameter ratio of the loading and the supporting ring, and is ~1/2 of the specimen thickness for a ratio of 1/2 and ~3 for a ratio

of 1/5 [¹³⁵,¹³⁶,¹³⁷]. In order to avoid non-linearity in the load-stress curve related with large deflection, the thickness has to be:

$$t \ge \sqrt{8\sigma \cdot r_1^2 / 3E}$$
 Eq. 29

Since the largest measured fracture stress was ~240 MPa, the calculated value $t \ge 230 \,\mu\text{m}$ was well below the thickness of the specimens (for strength measurement 0.5 – 2.8 mm). Specimen diameter, as well as thickness, also may affect the overall stress state. The stress state on the tensile surface inside the loading ring is, in theory, equibiaxial and constant. In practice, contact stresses are present under the loading ring. For round samples with D_s/t between 10 and 30, and $(D - D_s)/t$ between 1 and 6, (where *D* and D_s are diameters of the sample and support ring, respectively) the stresses within the loading ring are estimated to be constant within 2% [¹³⁸]. According to DIN 51105 [¹³⁹], the ratio D_L/D_s should be between 0.2 and 0.5 (where D_L is a diameter of loading ring). Taking the thickness range of the investigated spinel plates and the geometry of the bending set-up, following allowable ratios can be determined: $6.8 < D_s/t < 38$; $2.9 < (D - D_s)/t < 16$; $D_L/D_s = 0.5$. Both criteria are fulfilled (Fig. 18) for a thickness between 1.35 and 1.9.

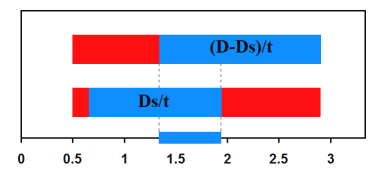


Fig. 18: The suitable thickness range of the specimens (in blue) for the ring-on-ring bending test.

The load-displacement curves were corrected for the compliance-related deformation of the set-up. To consider the machine compliance, a 25 mm-thick alumina specimen was subjected to loading using the configuration described above; up to the maximum force was used in the fracture tests of spinel specimens.

The actual machine deformation for a particular load was subtracted from the deformation obtained in the bending tests of the spinels in order to obtain an unbiased Young's modulus.

3.4 Indentation test

3.4.1 Macroindentation

Loads of 9.8, 49 and 98 N were applied for macrohardness determination (see sections 2.6.2, 4.4) by means of the hardness tester Buehler Met 10.

3.4.2 Depth-sensitive indentation

Experiments were carried out using both a Fischer HC100 indentation system and a combined nano-micro indentation set-up (CSM). Indentations were performed using loads between 0.1 and 20 N. Surface and cross-sections of the specimens were indented at low loads with the number of indents \sim 25, whereas for higher loads a sufficient statistical analysis could be obtained from 5 measurements. The available optical microscope system permits evaluation of the microstructural changes, as well as the crack lengths, for calculation of indentation fracture toughness (see section 2.6.3.1).

3.4.3 Hertzian indentation

Based on the Hertzian indentation method (see section 2.6.4.1), the local strength was estimated for the fine-grained material using spherical indenters with a radius of 0.5 and 2.5 mm, respectively. Impressions were made in the load range from 30 to 245 N, with load steps of 49 N. The critical load for the formation of cone cracks was estimated visually after the impression using an optical microscope.

3.5 Fracture toughness

3.5.1 Indentation

Using a macroindenter Buehler Met 10 and a microindenter Fischer HC100, the impressions were introduced into the spinel surface under different loads, ranging from 3 to 98 N (see section 2.6.3.1).

3.5.2 Indentation-Strength-Method

For ISM testing, typically up to four impressions were placed with an identical load (9.8 N) at the tensile specimen surface inside the loading ring area. This permits a clear determination of crack origin and an estimate of crack extension during the bending test before failure since fracture will typically only occur by crack extension starting from one of the pre-indentations. It was verified fractographically on every sample that failure in the subsequently carried out ring-on-ring bending test (see section 2.6.3.2) always occurred from one of the impression cracks.

3.5.3 Single edge notch bending

The measurements were carried out using an electromechanical testing machine (Instron 1362). The bending supports had an inner span of $S_2 = 7.5$ mm, and an outer span of $S_1 = 15$ mm. Due to limited availability of materials, only a small number of tests could be carried out with a specimen geometry of $4 \times 5 \times 25$ mm and $1.7 \times 3 \times 25$ mm for fine- and coarse-grained spinel specimens, respectively (see Table 5). Prior to the tests, the bar-shaped specimens were pre-notched following ASTM C1421 [¹⁴⁰].

3.6 In-situ fracture observation

In addition to fracture mode and crack shape estimation, which are critical parameters for fracture toughness determination (K_{IND}), it is necessary to know if indentation cracks and delaminations form during loading or unloading. Therefore, an indentation device with in-situ observation has been developed and applied (Fig. 19).

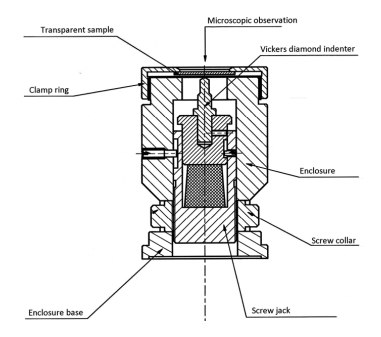


Fig. 19: Indentation device for in-situ fracture observation.

The specimen to be tested is fixed between a clamp ring and enclosure in order to avoid displacement of the specimen during loading. The gradual rotation of a screw collar moves the Vickers indenter to the contact surface, introducing the indentation mark and associated local fracture. A light microscope, Zeiss Axiomat, positioned above the specimen, permitted to observe the indentation process.

4 RESULTS AND DISCUSSION

4.1 Microstructure

The etched fine-grained and coarse-grained spinels were investigated via light microscope. Following the procedure outlined above (see section 3.2.2), the grain size was determined for both spinels, yielding an average grain size of ~ 5 μ m varying from 0.2 μ m to up to 50 μ m for fine-grained transparent spinel (Fig. 20a), and an average grain size 60 μ m, with a size variety from 5 μ m up to 170 μ m for the coarse-grained transparent spinel. The microstructure of both spinels is inhomogeneous, and partially consists of zones with relatively coarse and fine grains (Fig. 20).

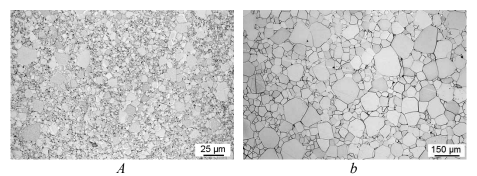


Fig. 20: Thermally etched structures of fine-grained (a) and coarse-grained (b) spinel.

In addition, a thin lamella was extracted by FIB from the fine-grained material and then characterized by TEM, confirming the existence of nanometer-sized grains and well-defined grain boundaries (Fig. 21a). A complementary EDX element analysis verified that the material consists locally only of aluminum, magnesium and oxygen (Fig. 21b). The presence of copper and carbon in the spectrum is caused by the material of the specimen holder and the carbon coating deposited to enhance electrical conductivity, respectively.

TEM investigation was applied for analyzing the element composition between fine- and coarse-grained areas in fine-grained spinel.

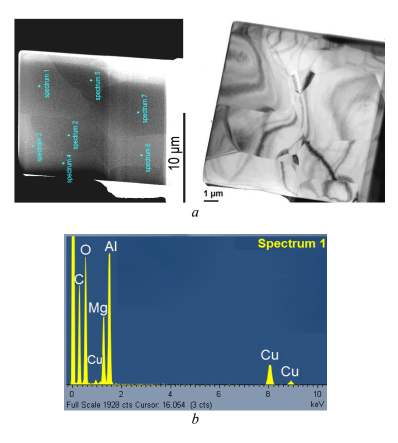


Fig. 21: Fine-grained spinel lamella investigated via (a) TEM and (b) EDX.

4.2 X-ray analysis

An X-ray phase analysis confirmed that there are no evident impurities in the fine-grained and coarse-grained transparent spinels, proving *fcc* cubic spinel crystal structure (Fig. 22, Fig. 23). The obtained lattice parameter, 0.808 nm, coincides with literature findings (see section 2.1). Therefore, XRD and EDX element analyses

confirmed the purity of the magnesium aluminate spinel, permitting the comparison of the mechanical data to the literature values of this material.

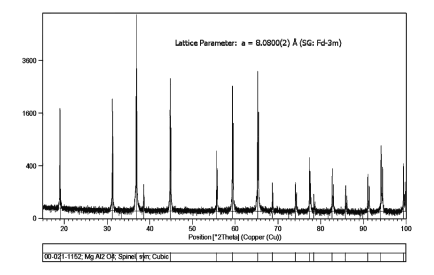


Fig. 22: XRD diagram of the fine-grained transparent magnesium aluminate spinel

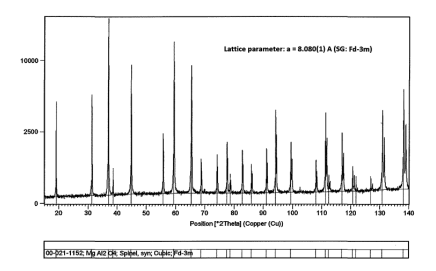


Fig. 23: XRD diagram of the coarse-grained transparent magnesium aluminate spinel

4.3 Young's modulus

Young's modulus was determined using bending, impulse excitation and indentation testing (see sections 2.6.1.1, 2.6.1.2, 2.6.1.3). The value for fine-grained spinel derived from the ring-on-ring bending tests without compliance correction decreases significantly, with increasing specimen thickness from 270 GPa down to ~ 160 GPa (~ 45 %) (Fig. 24).

The circle-shaped data points in Fig. 25 represent Young's Moduli from ring-onring bending tests after compliance correction, as outlined in section 3.3.2. The deformation of the experimental setup used for compliance calibration (see section 3.3.2) was ~3.5 μ m in the load range 550 - 650 N, whereas for fine-grained spinel the deflection of the specimens was ~20 μ m at the same load range. Assuming an uncertainty in the compliance calibration of 1 μ m leads to an uncertainty in the experimentally determined modulus of ~7% for fine-grained spinel. The value measured for the thinnest specimens is ~270 GPa, whereas for larger thicknesses the modulus decreases by ~10% (Fig. 25). Therefore, this effect is close to the limits of uncertainty.

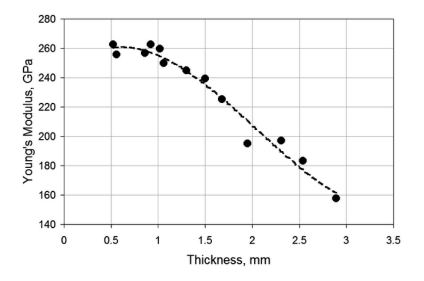


Fig. 24: Apparent Young's modulus (not corrected for compliance) of fine-grained spinel as a function of the specimen thickness.

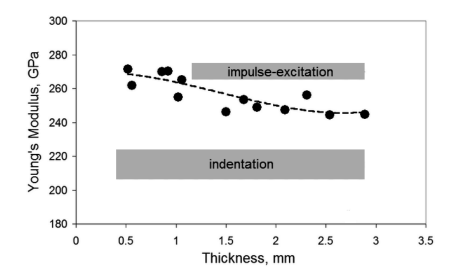


Fig. 25: Apparent Young's modulus as a function of specimen thickness. Results are derived from compliance-corrected load-displacement curves obtained in ring-on-ring bending tests for fine-grained spinel specimens.

The value of ~270 GPa determined for thin specimens (~1 mm) coincides with the average modulus determined using the impulse excitation technique (~270 \pm 5 GPa), whereas the depth-sensitive indentation technique yielded a 20% lower value (~210 \pm 10 GPa, for loads from 0.5 to 1 N) (Fig. 25). The low Young's Modulus obtained by indentation can be explained by surface effects [¹⁴¹].

The same bending test procedure was also used for the coarse-grained spinel. However, specimens were only available with a thickness of around 2 mm. After compliance correction, the ring-on-ring bending tests yielded E_{ROR} values of 242 ± 13 GPa, whereas impulse excitation and indentation tests resulted in Young's moduli of ~270 ± 5 and ~225 ± 10 GPa, respectively.

Overall, the data for fine- and coarse-grained material coincide only for impulse excitation. The bending tests yield lower values, which might be biased by morphological effects, e.g., the grain size and local defects (Fig. 20, Fig. 54). Indentation tests reveal \sim 5% higher values for the coarse-grained spinel, compared to the fine-grained material. This difference might be related to the number and amount of the coarse-grained areas with higher local Young's modulus in the fine-grained material (see section 4.4). The

nearly identical value of \sim 270 GPa determined by impulse excitation technique for both spinel variants reflects the low sensitivity of this technique to local structural inhomogeneities. The result obtained using this method represents the integral property of the entire specimen volume.

The local indentations into grains (see section 4.4) were analyzed in more detail to obtain insight into the origins of the differences between indentation moduli and the global values obtained from bending and impulse excitation tests. The microindentation into the thermally-etched grain structure (see section 3.1.3) verified that (for fine-grained spinel in particular), the modulus is also affected by the local structure of the material. For instance, fine-structured spinel areas (grain size $0.5 - 5 \mu m$) lead to a large scatter from 140 to 240 GPa, whereas coarse-structured spinel areas in the fine-grained material (grain size $20 - 60 \mu m$) revealed data with less deviation (from 190 - 230 GPa). Although thermal etching appeared to increase the average Young's modulus by ~10% (\sim 5% for the hardness), it can be concluded that the average of the local properties is not equal to the global properties determined by bending and impulse excitation, due to the different integral effect of grain boundaries and defects. A similar behavior could not be confirmed for the coarse-grained spinel (see section 4.4). Consequently, these data revealed less scatter in comparison with fine-grained spinel, due to different morphology of the material (big single grains and grain boundaries of relatively fine grains between them yield ranges of 170 – 190 GPa and 210 – 230 GPa, respectively).

In addition, the temperature dependency of the Young's modulus (up to 1000 °C) was determined for the fine- and coarse-grained spinels using bar-shaped specimens by impulse-excitation testing in air. Plates could not be tested at high temperature using the available experimental setup. The temperature dependency corresponds with the decrease of 1 % per 100 K quoted in literature for ceramic materials [¹⁰⁶] (Fig. 26). Since the temperature dependence of the Young's modulus is an intrinsic materials behavior, it might considered to be representative also for the coarse-grained batch. Indeed, for both spinels Young's modulus linearly decreases with temperature increasing. Only one measurement was carried out for each of both spinels and the respective Young's modulus shows a difference of ~ 5 % over the entire temperature interval. However,

Young's modulus of the coarse-grained spinel decreases at a slightly stronger rate with increasing temperature.

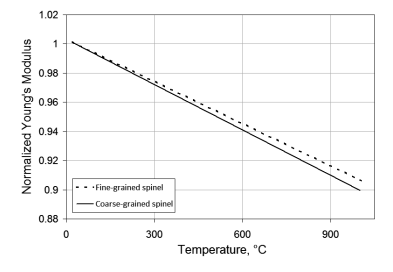


Fig. 26: Normalized Young's Modulus of the fine- and coarse-grained spinels as a function of temperature determined by impulse excitation (E_{IE}) .

4.4 Hardness

Impression tests were carried out for a load range from 1 to 98 N in the specimen surface using micro- and macro-indentation (see sections 3.4.1, 3.4.2). Three impressions were applied at loads of 1, 3, 9.8, 49 and 98 N each. Fig. 27 illustrates the load-dependency of hardness, which is less pronounced for the fine-grained spinel. The hardness becomes almost load-independent above 1 N for the fine-grained spinel, whereas for the coarse-grained spinel the hardness is still load-dependent even at 96 N.

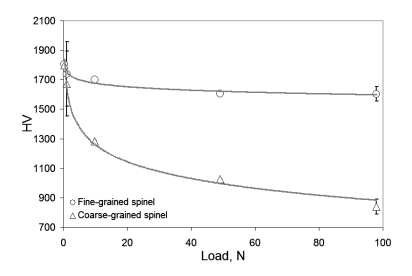


Fig. 27: Vickers hardness - indentation load dependency of the fine-grained and coarsegrained spinel (surface).

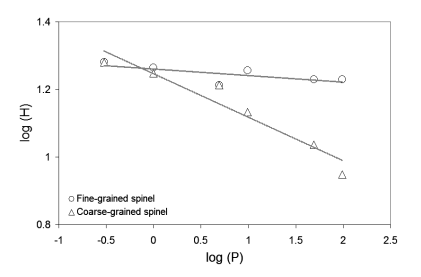


Fig. 28: Indentation size effect for fine-grained and coarse-grained spinel materials.

Such load dependencies of indentation hardness were often reported for ceramics [e.g. ¹⁴²] and are usually referred to as indentation size effects. A clearer representation of

the effect can be obtained from a *log-log* plot of the data (Fig. 28), yielding rather linear dependencies. A larger slope is obtained for the coarse-grained material. Literature has suggested that a larger slope indicates that the material that is less prone to impact damage [143 , 144].

Note that a load of 1 N yielded an average indentation hardness of 16 ± 2 GPa for both materials for impressions carried out in surface and cross-section.

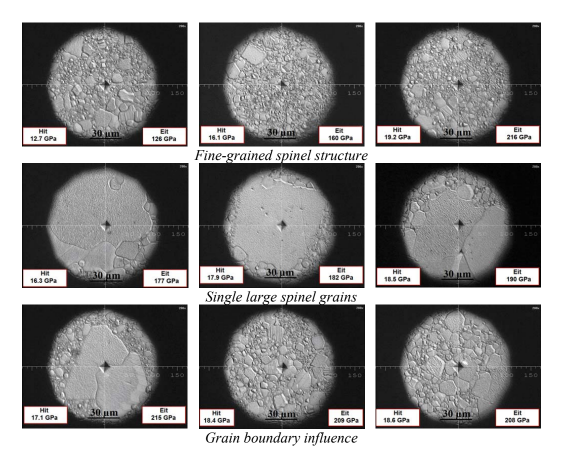


Fig. 29: Typical hardness impressions in the fine grained spinel surface (P = 1 N)

Additional indentation tests were carried out to determine if a correlation of microstructure and local hardness exists. Before indentation, the fine-grained and coarsegrained specimens were thermally etched as described in the experimental section. The negligible increase of the average hardness of 5% after etching verifies that the etched specimens are comparable to the initial condition.

As indicated in Fig. 29, impressions in the larger grains $(20 - 60 \ \mu\text{m})$ yield hardness values of 16 - 18 GPa (P = 1 N). The impressions into nano/micro-grained spinel areas lead to larger hardness variations, with values in the range of 13 to 19 GPa that might be attributed to the effect of structural inhomogeneities, such as grain boundaries, twin boundaries, inclusions or precipitates.

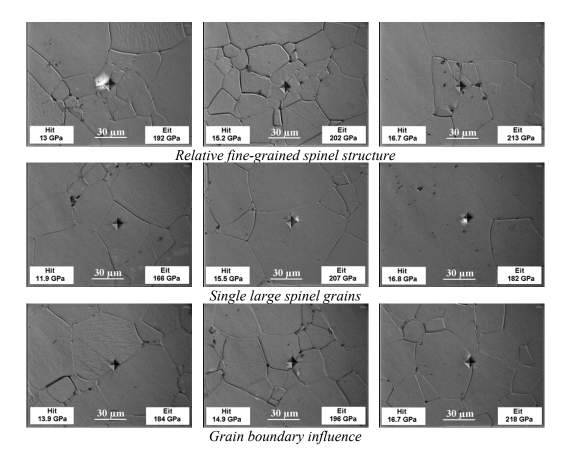


Fig. 30: Typical hardness impressions in the coarse-grained spinel surface (P = 1 N)

For the coarse-grained material, a hardness range from ~ 12 to 17 GPa was obtained for all local grain sizes (Fig. 30). Overall, the data do not lead to conclusions regarding grain size effect.

4.5 Fracture toughness

4.5.1 Indentation

The calculation of indentation fracture toughness requires first a determination of the crack shape. An accurate way to determine the indentation crack shape is to break up a specimen after applying a line of indents (here load 49 N, distance between indents ~500 μ m). After manual breakage, it could be confirmed that the cracks have a half-penny shape for both spinels (Fig. 31), an important prerequisite for choosing the relevant geometric factor in the fracture toughness relationship (see section 2.6.3.4), which was here (Y/Z) ~ 1.27.

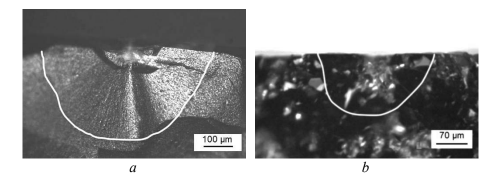


Fig. 31: Half-penny shaped crack starting at a Vickers indent (P = 49 N): a) finegrained and b) coarse-grained spinel.

However, crack growth appears also to be influenced by the local stress field. In indentation testing, the radial cracks in fine-grained spinel are formed in a mixed transand inter-granular fracture mode (Fig. 32a), whereas the central region beneath the Vickers impression (which is under compressive stress), reveals an intergranular fracture (crushing) (Fig. 32b). Since for coarse-grained spinel the use of higher loads lead to chipping of the indented areas, whereas no radial cracks could be obtained at low loads, the fracture mode could not be characterized for this material. But it does not appear farfetched to assume that the transgranular fracture prevails during the indentation for both regions of the impression.

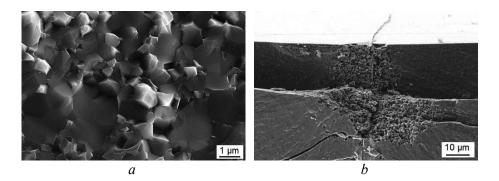


Fig. 32: Fracture path in the fine-grained material *a*) after bending test ($\sigma = 135$ MPa) and *b*) central region under the indentation (P = 49 N).

The device, developed in-house and built to observe hardness impression tests insitu (see section 3.6), allowed further analysis of indentation crack formation in transparent ceramics. The fine-grained spinels revealed radial crack formation (Fig. 33a) during loading, whereas a test carried out for comparative purposes at silica glass showed cone crack formation (Fig. 33b).

Despite the clear images obtained using the optical microscope, the very narrow crack tips are not clearly visible in reflecting light images (Fig. 34a). In Fig. 34b, the same impression depicted via transmitted light microscope shows a sharper contour and the length of the lateral, as well as of the radial cracks, which can be measured more accurately. The difference between the results obtained using reflected and transmitted light is ~ 10 %. Hence, the use of transmitted light is recommended.

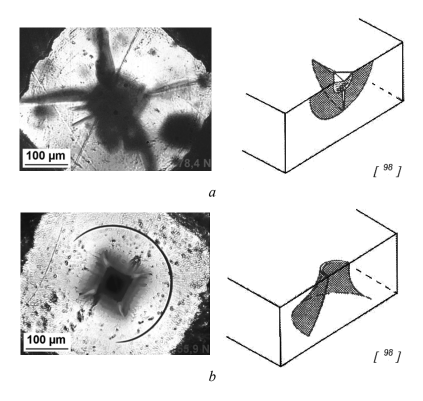


Fig. 33: In-situ fracture observation and schematic crack morphology at indentations in a) fine-grained spinel and b) silica glass.

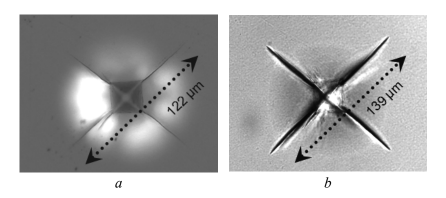


Fig. 34: Vickers impression in fine-grained spinel (P = 9.8 N), analyzed using a) reflected and b) transmitted light microscopy.

To determine the indentation fracture toughness, the relationships by Lankford et al. [99], Niihara et al. [93] and Anstis et al. [94] were selected in accordance with the observed crack shape (see section 2.6.3.1). The obtained fracture toughness values remained unchained, independent of the applied loads in the load range from 3 to 98 N for the fine-grained spinel (Fig. 35 and Table 7). However, high loads lead to large delamination and spallation for coarse-grained spinel (Fig. 38b). Therefore, all further investigations were performed with a load of 3 N (Fig. 36), which resulted in well-defined radial cracks (Table 7).

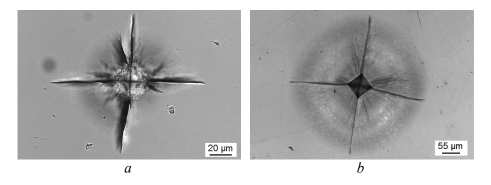


Fig. 35: Vickers impressions in fine-grained spinel for loads of a) 9.8 N and b) 49 N.

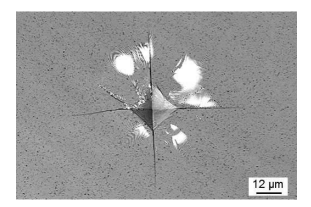


Fig. 36: Vickers impression in etched coarse-grained spinel for a load of 3 N.

	<i>K_{IND}</i> ,(MPa·m ^{0.5}) for fine-grained spinel	<i>K_{IND}</i> ,(MPa·m ^{0.5}) for coarse-grained spinel
<i>K_{IND}</i> (<i>Lankford</i> et al. [⁹⁹], c/a > 2)	ime-gramed spiner	coarse-gramed spiner
loads 3 N		2.2
5 N	2	
9.8 N	2.1	
49 N	1.9	
98 N	2.1	
<i>K_{IND} (Niihara et al. [⁹³], c/a</i> > 2.5)		
loads 3 N		1.2
5 N	2	
9.8 N	1.9	
49 N	1.8	
98 N	2	
K_{IND} (Anstis et al. [⁹⁴], $c \ge 2a$)		
loads 3 N		0.8
5 N	1.2	
9.8 N	1.3	
49 N	1.2	
98 N	1.3	

Table 7. Indentation fracture toughness (K_{IND}) as a function of load

* for all obtained values the uncertainty is ~ 0.05

The K_{IND} values for fine-grained spinel reveal a slight difference of ~5% between the results obtained using the equation by Lankford et al. [⁹⁹] and Niihara et al. [⁹³], whereas in case of the coarse-grained spinel the difference is ~45%. The calculated K_{IND} value based on the equation by Anstis et al. [⁹⁴] results in the lowest fracture toughness value of ~1.3 MPa·m^{0.5} for the fine- and ~0.8 MPa·m^{0.5} for the coarse-grained spinel. Note that all equations consider a half-penny shaped crack. The difference between these equations is the adjustment of a calibrated geometrical factor based on industrial ceramics experiments (see section 2.6.3.1) [^{94, 145}]. Based on the results obtained in the current study using ISM and SENB methods, as well as the Griffith criterion (see sections 2.6.3.2, 2.6.3.3, 2.6.3.4), it is suggested that the relationships suggested by Lankford et al. [⁹⁹] and Niihara et al. [⁹³] are most appropriate for evaluating the fracture toughness of the fine-grained spinel material currently studied.

4.5.2 Indentation-Strength-Method

Fracture loads obtained in ring-on-ring bending tests for pre-indented specimens (ISM, see sections 2.6.3.2, 3.5.2) [⁴⁰] were also used to determine the fracture toughness (K_{ISM}) of coarse- and fine-grained spinels. Before the ISM test, typically up to four impressions are placed at the tensile loaded ring-on-ring bending specimen surface. It was verified fractographically for every tested specimen that failure occurred from only one of the impression cracks. The reduction of average fracture stress due to the pre-indentation cracks of $c_0 = 67 \,\mu\text{m}$ was ~35% compared to material without indentation cracks, verifying again that indentation cracks were the failure origins in these bending tests. The indentation cracks grow subcritically during the bending test, until one of them results in fracture (Fig. 37). The validity of the ISM also depends on the subcritical crack extension before failure, since this parameter enters the calculation via the calibration factor A (Eq. 9).

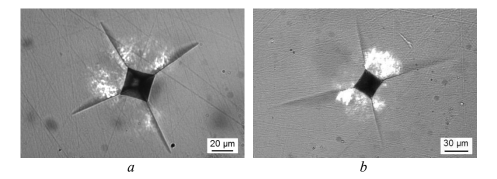


Fig. 37: Indentation cracks in fine-grained spinel after indentation loading 9.8 N. a) before and b) after bending test

Therefore, the indentation crack size was measured before and after the test, and the crack length after stable crack growth c_m was estimated from the three impressions that did not lead to failure. This yielded a value of $130 \pm 10 \ \mu\text{m}$, which was twice the

initial crack length. Since the calibration factor is proportional to the ratio of the crack length to the power of (3/8), a 16% larger *A* factor had to be used in Eq. 9. Finally, a value of $K_{ISM} \sim 1.9$ MPa·m^{0.5} was obtained, which is similar to the K_{IND} results, further verifying the methodology used. However, the use of the ISM test for coarse-grained spinel was not possible, since starting from 5 N the radial cracks and the associated intergranular fracture lead to delamination of grains (Fig. 38). Note that defined crack extension during bending is one of the main conditions, which must be satisfied for the ISM.

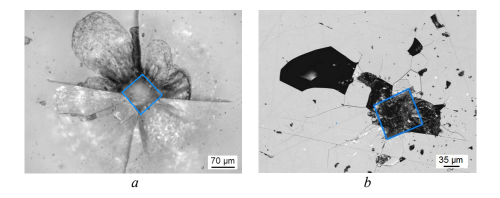


Fig. 38: Vickers impressions (P = 49 N) into a) fine-grained spinel and b) coarsegrained spinel (P = 49 N).

4.5.3 Single edge notch bending

The K_{SENB} was determined for fine- and coarse-grained spinels from four-point bending tests (see section 2.6.3.3) following ASTM C 1421. The fine-grained spinel shows higher single edge notch bending toughness 1.8 ± 0.2 MPa· \sqrt{m} compared with coarse-grained spinel 1 ± 0.4 MPa· \sqrt{m} . The lower K_{SENB} of coarse-grained spinel and its higher scatter might be partially attributed to the existence of weak grain boundaries (inherent defects), (see section 4.8), but also the limited number of available specimens (see section 3.1.2).

4.5.4 Griffith criterion

This method requires fractographic analysis of the broken specimens and localization of the fracture origin. The size of the defect originating the crack and the fracture stress determined in the ring-on-ring bending test can then be used to calculate fracture toughness using Eq. 12. Fractography yielded defect sizes from 200 μ m to almost 1 mm (Fig. 51, Fig. 53), yielding fracture toughness values of ~ 1.6 ± 0.2 MPa· \sqrt{m} , coinciding with the data obtained by the other two methods (see Table 8).

Method	Fine-grained spinel	Coarse-grained spinel	
	Lankford		
	2 ± 0.1	2.2 ± 0.1	
<i>K_{IND}</i> , MPa·√m	Niihara		
	1.9 ± 0.1	1.2 ± 0.05	
	Anstis		
	1.3 ± 0.05	0.8 ± 0.05	
K_{ISM} , MPa· \sqrt{m}	1.9 ± 0.2		
K_{SENB} , MPa· \sqrt{m}	1.8 ± 0.2	1 ± 0.4	
<i>K_{IC} (Griffith)</i> , MPa·√m	1.6 ± 0.2		

Table 8. Fracture toughness obtained by different methods

The SENB test suggests that in the case of indentation, the Niihara equation [⁹³] is the most appropriate for estimating the fracture toughness of both spinels.

4.6 Photoelasticity

The photoelastic effect (see section 2.6.6) can be used for a qualitative analysis of local strains that might be associated with structural inhomogeneities and defects [¹³⁰,¹³¹]. The strain field induced by Vickers indentations and secondary cracks in a spinel fragment after bending was studied. Observation via a polarization microscope revealed

two main areas of higher light intensities, one close to the center of the impression and at the end of a radial crack tip, and the other at the end of a secondary crack tip that remained after bending tests (Fig. 39a, b), respectively. The light intensity close to the center of the impression could be related to lateral cracking and associated interference effects, and therefore was an artifact unrelated to residual strains. However, closer observation also revealed higher intensities near the radial crack tips, induced by indentation, which coincided with observations made at secondary crack tips in the fractured bending specimens.

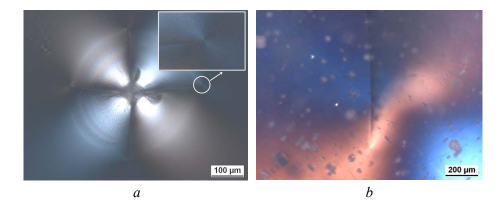


Fig. 39: Light intensities of strained zones: a) at crack tips (marked white) initiated by indentation (P = 49 N); b) at a crack in a spinel fragment after bending ($\sigma = 147$ MPa).

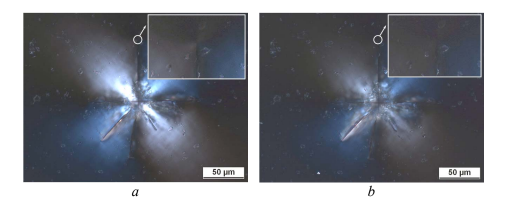


Fig. 40: Strained zones at one impression (P = 9.8 N) a) before and b) after heat treatment.

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Subsequent annealing at (1000 °C, 1h) in air appeared to relax deformations and strains, particularly the size of the local strain field surrounding the crack tip (Fig. 40).

4.7 Fracture strength

In initial experiments, the effect of polishing conditions on fracture strength of fine-grained spinel was tested for three different surface states (see section 3.1.2) in order to optimize the preparation process. The Young's Modulus was, as expected, independent of the polishing procedure, whereas the strength increased (from ~ 130 MPa up to ~ 160 MPa) when the roughness was decreased from low to medium quality, and remained constant at high-quality polishing. The lower Weibull modulus of the high quality polished batch can be explained by remaining local scratches after machining. As expected, the surface condition has an influence on the fracture strength for large roughness values (Table 9), and therefore the average polishing procedure was used for further experimental work (see section 3.1.2).

Roughness, R _a	σ _f , MPa	<i>E_{ROR}</i> , GPa	т
2.34	132 ± 21	233 ± 6	8 ± 2
1.3	165 ± 32	240 ± 8	6 ± 2
0.83 - 1.26	162 ± 46	234 ± 6	4 ± 1

Table 9. Effect of three different polishing qualities on the mechanical properties of finegrained spinel

The fracture stress measurements yielded a fracture strength of $\sigma_f = 155 \pm 38$ MPa for all fine-grained spinel batches of average and high quality polishing (~150 specimens), which is slightly lower than the values determined for individual batches of ~30 specimens (see Table 9). Contrary to the Young's modulus (see section 4.3), the fracture stresses did not show a dependency on specimen thickness (1.3 – 2.8 mm) within the limits of experimental uncertainty. Hence, in agreement with ASTM C1499, the fracture stress and therefore strength, is less sensitive to thickness variations. From the Weibull representation of the results (Fig. 41), a characteristic strength of $\sigma_0 = 169 \pm 3$ MPa

was derived for fine-grained spinel. The Weibull modulus reflecting the scatter of the results is rather low with $m \sim 5 \pm 2$. The confidence interval is 4.4 < 5 < 5.5 (see section 2.6.4).

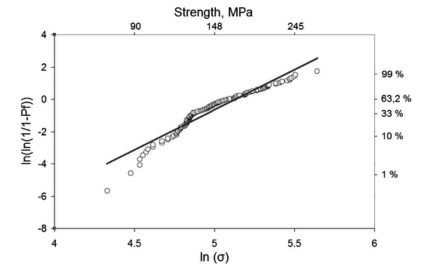


Fig. 41: Two-parameter Weibull plot of the fracture stress data of fine-grained spinel.

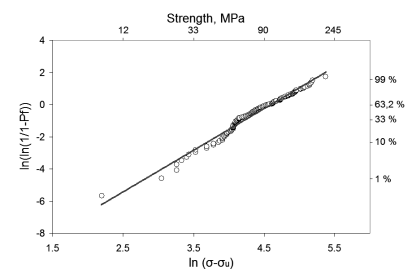


Fig. 42: Three-parameter Weibull plot of the fracture stress data for the fine-grained spinel.

Utilizing a three-parameter Weibull distribution leads to a better description of the data and therefore to a higher correlation coefficient R^2 (~0.92 for the two- and ~0.98 for the three-parameter Weibull distribution). The threshold stress σ_u was obtained by optimization of R^2 to $\sigma_u = 67$ MPa. The three-parameter Weibull distribution yields a characteristic strength $\sigma_0 = 166 \pm 3$ MPa for fine-grained spinel. The lower Weibull modulus of $m = 2.6 \pm 0.3$ (Fig. 42) is typical for the description of data with a three-parameter distribution [¹⁴⁶].

Due to the limited number of ~30 specimens the data obtained for the coarsegrained materials were only described by a two-parameter distribution since an amount of ~100 specimens is required for three-parameter Weibull distribution [¹⁴⁷,¹⁴⁸]. The average fracture stress is $\sigma_f = 73 \pm 9$ MPa, the characteristic strength $\sigma_0 = 77 \pm 1$ MPa and the Weibull modulus $m = 10 \pm 2$ (Fig. 43). Therefore, the coarse-grained spinel has a lower strength, but a higher Weibull modulus than the fine-grained material. From this, it can be concluded that defects in the coarse-grained spinel are larger but distributed more uniformly. For coarse-grained spinel, the confidence interval is 7.7 < 10.2 < 12.4.

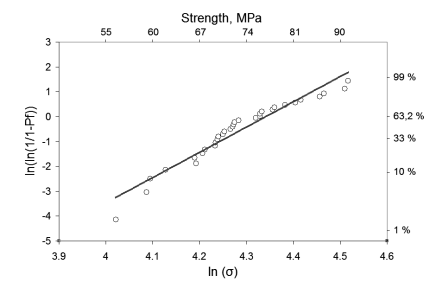


Fig. 43: Two-parameter Weibull plot of the fracture stress data of the coarse-grained spinel

4.7.1 Weibull modulus uncertainties

The accuracy of statistical evaluation mostly depends on the number of available specimens N and the Weibull modulus m. In addition, a program compilation via JavaScript software permitted to partially evaluate the available data sets.

A procedure assesses the effect of dataset selection on the uncertainty of statistical strength evaluation of the fine-grained spinel. From the entire set, 30 fracture stress values were selected 10 times randomly, and the maximum and minimum Weibull modulus was calculated. The number of selections of the 30 data values was increased in subsequent calculations to 30, 60 and 90, in order to determine whether there is any effect on the resulting maximum and minimum values. The range of uncertainty of the Weibull modulus increases from 4.2 < m < 6.3 for 10 loops to 4 < m < 6.7 for 90 loops (Fig. 44). This compares quite well to the standard deviation of s (m = 5) = 1 and confidence interval of 4.4 < 5 < 5.5 (see section 2.6.4). The influence of the number of cycles on the Weibull modulus seems to be negligible, since the Weibull modulus only was changed by $\sim 5\%$.

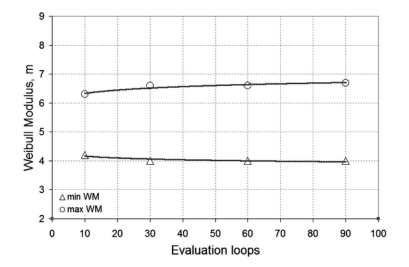


Fig. 44: Relation between number of calculation cycles and obtained Weibull moduli

4.7.2 Particle-size distribution

Fine-grained spinel contained relatively large agglomerates that affected the transparency. As a result, one question to be answered concerned whether these agglomerates also affect the strength (Fig. 45, Fig. 46).

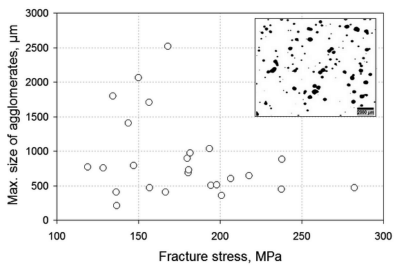


Fig. 45: Fracture stress as a function of the maximum size of agglomerates.

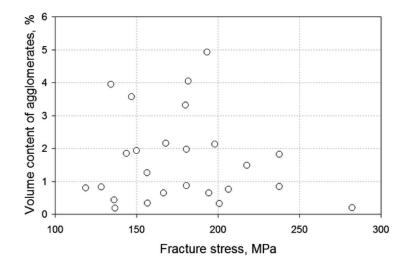


Fig. 46: Fracture stress as a function of the specific volume of agglomerates.

Maximum size and volume of agglomerates were provided for selected finegrained specimens by the project partner CeramTec-ETEC, as obtained by light microscopy and subsequent quantitative image analysis (see section 3.2.2). Maximum size and volume of agglomerates for the characterized 25 fine-grained spinel specimens were within the interval $210 - 2530 \mu m$ and 0.17 - 4.9 %, respectively. The specimens were fractured in ring-on-ring bending tests. Fig. 45 and Fig. 46, which display the fracture stress values versus maximum defect size and volume, respectively, illustrate that there is no correlation of fracture stress with either of these parameters. As a result, such visible defects are not failure-relevant for this material.

4.7.3 Local strain field

Although a macromechanical effect of optically visible agglomerates can be ruled out, the micromechanical effect still requires assessment. Therefore, in an extension of the photoelastic analysis of the strain at hardness impressions (see section 4.6), the photoelastic method was also used to qualitatively determine the local strain field at optically visible agglomerates. Fig. 47 clearly verifies the existence of a residual strain field around these agglomerates. After an additional heat treatment T = 1000 °C for 1 h (Fig. 47b) (see section 3.1.3), the local strains appear to be slightly lower due to relaxation. In an attempt to assess whether heat treatment affects fracture stress, three specimens were subsequently fractured in ring-on-ring bending tests. The average fracture stress 137 ± 9 MPa and Young's modulus of 235 ± 5 GPa were obtained for heat treated fine-grained specimens, which corresponds with the as-received state. However, the Young's modulus determined using indentation (24 indents in the surface) increased from 205 ± 12 to 232 ± 9 GPa, and hardness from 17.2 ± 1 to 17.7 ± 1 GPa compared to the same specimen in the as-received state. This slight effect, especially on the Young's modulus, might be related to oxygen diffusion or reactions in the surface areas that do not affect the global properties obtained in a bending test $[^{149}, ^{150}]$.

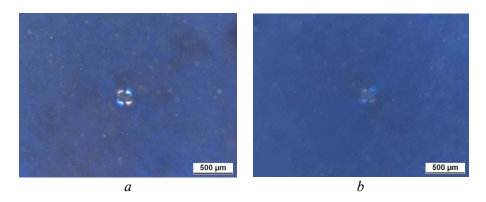


Fig. 47: Photoelastically determined residual stress field around agglomerates a) before and b) after heat-treatment.

Also tested was whether the local agglomerates influence crack growth. A crosssection through a coarse-grained spinel agglomeration was loaded by a wedge after an additional notch had been introduced. Polarization microscopy revealed that the agglomerate was surrounded by a local residual strain field (Fig. 48a). In the subsequent wedge test, the crack bypassed the agglomerate shown in the SEM image in Fig. 48b. Even subsequently introduced indentations did not lead to a fracture path through the particle, suggesting the existence of a local compressive strain field surrounding the particle. Therefore, even though a direct correlation of the macromechanical properties with the agglomerates was not possible, the micromechanical tests revealed that crack growth is affected by these inhomogeneities of microstructure and in fact, suggests that the optically visible agglomerates enhance the micromechanical properties. The absence of any effect in the macromechanical test might be related to the limited number of specimens, or that the majority of the optically visible agglomerates are not related to local strain fields.

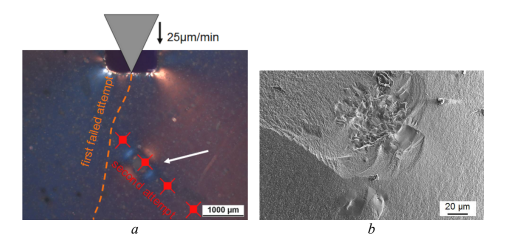


Fig. 48: Influence of a coarse-grained agglomerate on fracture path. a) Sketch of the implemented measurements (the agglomerate is indicated by an arrow); b) SEM image of the agglomerate after indentation.

4.8 Fractography

After ring-on-ring bending testing, specimens were investigated with respect to their fracture modes. As shown in Fig. 49a, a mixture of inter- and transgranular fracture paths is typical for fine-grained spinels. A transgranular fracture path is visible in local structures with grain sizes larger than 1 μ m, whereas intergranular fracture was observed in the nano-grained regions (0.2 – 1 μ m). The coarse-grained spinel failed only in an intergranular fracture mode (Fig. 49b).

However, the stresses field at large-grained agglomerates in fine-grained spinel also appears to influence the fracture mode. Indentation testing leads to radial cracks in a mixed mode, whereas the central region beneath a Vickers impression (which mainly undergoes compressive stresses) reveals intergranular fracture (crushing) (Fig. 32b).

In general, the number of fractured pieces in the biaxial ring-on-ring tests appeared to scale with the fracture stress (Fig. 50); i.e., more pieces were obtained for a specimen that fractured at high stress. This study observed similar fracture patterns to ones reported by Quinn [151]. Characteristic positions (kinks and crack undulations) in

selected specimens that failed at low fracture stresses (corresponding to the largest defects) were chosen for subsequent fractographic analysis.

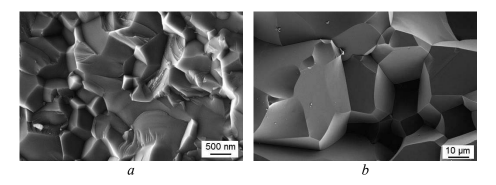


Fig. 49: a) Mixed trans- and inter-granular fracture in fine-grained spinel and b) intergranular fracture in coarse-grained spinel.

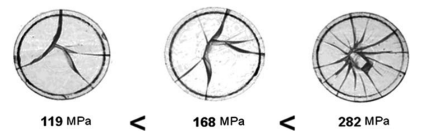


Fig. 50: Specimens after fracture tests illustrating the correlation of fracture stress and number of fractured pieces. The measured fracture stress values are indicated.

Optical micrographs and SEM images suggest that coarse-grained agglomerates and associated local microcracks are the failure origin for the fine-grained material (Fig. 53). In fact, these results confirm a strong influence of surface preparation and surface condition (see sections 3.1.2 and 4.7) on strength, since crack-like features surrounding the large grains are not observed in fracture surfaces. Hence, the inherent local microcracks in the vicinity of coarse-grained agglomerates might have already existed in the raw material and then extended during the grinding and polishing. The correlation of

individual fracture stresses with defect sizes was also used to estimate fracture toughness (see section 4.5.4). Initial defect sizes measured at fracture surfaces ranged from 150 μ m up to ~1 mm (Fig. 51, Fig. 52, Fig. 53). This correlation is illustrated in Fig. 51, which shows an agglomerate of large grains (size ~ 1 mm) and associated microcracks before and after a bending test, and clearly identifies this agglomerate as failure origin.

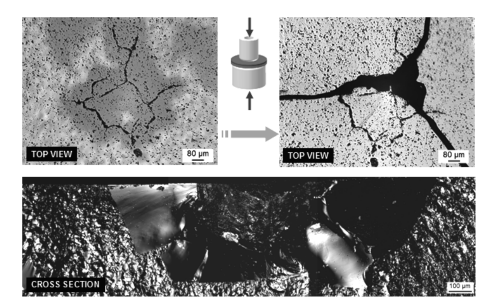


Fig. 51: A typical surface defect as fracture origin in a fine-grained spinel specimen. a) agglomerate and associated microcrack before the test, b) reassembled specimen after the test, confirming the failure origin and c) fracture surface confirming the extent of the coarse-grained zone.

SEM/EDX analysis of a similar coarse-grained area verified that the chemical composition of fine-grained structures and coarse-grained agglomerates are identical (Fig. 52b).

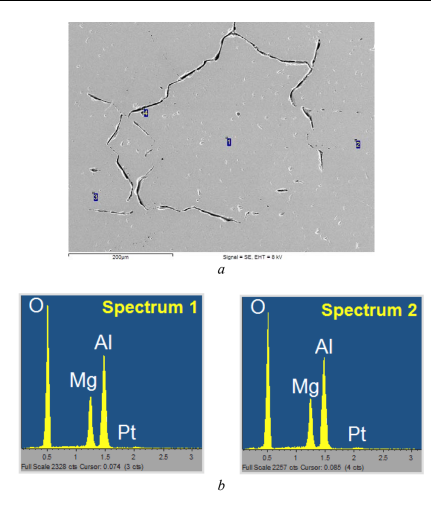


Fig. 52: SEM/EDX analysis of fine-grained structures and coarse-grained agglomerates.

Other typical failure origins in fine-grained spinel specimens, shown in Fig. 53, confirm the correlation of crack origin and coarse-grained agglomerates. Typical fracture originating agglomerates with sizes from 200 to 450 μ m have been detected. Note that overall the agglomerates appear to be related to failure, and act as failure origin. Optically visible agglomerates do not appear to be responsible for failure, suggesting that different kinds of agglomerates exist in the material. Furthermore, micromechanical tests suggest a compressive stress field around the agglomerates, which should be balanced by tensile

stresses in the agglomerates – another possible reason for microcracks, which could be initiated and enhanced by polishing.

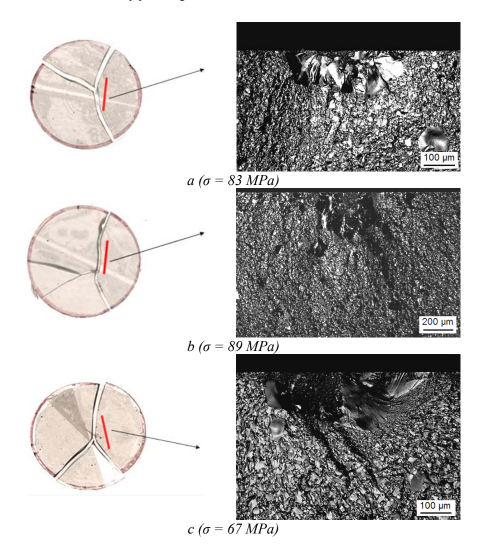


Fig. 53: Fracture origin determination after ring-on-ring bending tests.

Weak grain boundaries and pores appear to be more pronounced for coarsegrained spinel, and seem to be characteristic crack initiating defects (Fig. 54). The length of weak boundaries can reach 200 μ m and therefore might be considered as fracture origins.

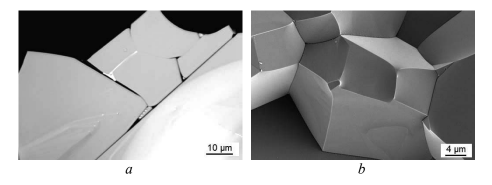


Fig. 54: Inherent porosity (a) and weak grain boundaries (b) as possible fracture origins for the coarse-grained spinel.

4.9 Statistical analysis

4.9.1 Area effect on fracture strength

The area effect on fracture strength was calculated (Fig. 55) using statistical analysis (see section 2.6.5) in order to estimate the properties of life-size components. The representations are based on data from ring-on-ring bending (characteristic strength $\sigma_0 = 169$ and 77 MPa for fine-grained and coarse-grained spinel, respectively). The figure illustrates how strength decreases with increasing area at a given thickness of ~2 mm. Note that the coarse-grained spinel is less sensitive to the area effect due to its higher Weibull modulus (see section 4.7), but generally yields a lower strength.

Based on the Hertzian theory (see section 2.6.4.1), the local strength was estimated for the fine-grained material using two spherical indenters of different radii (0.5 and 2.5 mm, respectively). Impressions were made in the load range from 30 to 245 N, with load steps of 49 N. The critical load for the formation of cone cracks was estimated visually after the impression, using an optical microscope. The estimated critical load P_c was used to calculate contact areas by Eq. 24 and local strengths by Eq. 23. The results for the fine-grained spinel are given in Table 10 and depicted in Fig. 56.

Although the estimate of critical load includes a rather large uncertainty of ~40% due to the discrete loading steps 49 N of the used machine, it can be seen that the data agree with an extrapolation of the results from ring-on-ring bending tests to lower deformed areas (see Fig. 56).

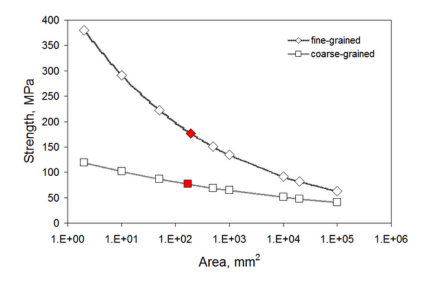


Fig. 55: Area effect on fracture strength for fine- and coarse-grained spinels. The experimental strength value is marked by red color.

Table 10. Obtained results for fine-grained spinel after Brinell indentation.

<i>R</i> , mm	P_c , N	$A_{eff.Hertzian},\mathrm{mm}^2$	$\sigma_{Hertzian},{ m MPa}$
0.5	98	1.4	470
2.5	196	6.6	287

The effective area deformed under tensile stress in bending was calculated according to Eq. 19 and according to Eq. 24 for spherical indentation. The stress gradient in the surface has been considered in these equations.

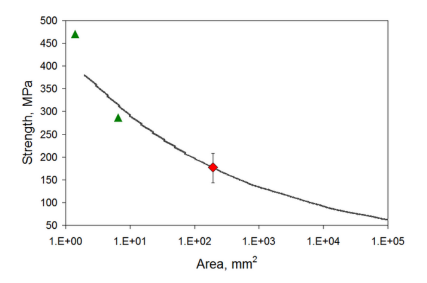


Fig. 56: Area effect on fracture strength in bending (black trend calculated from the experimental value (red) and Eq. 20) and from Brinell indentation (green triangles) for fine-grained spinel.

4.9.2 Slow crack growth behavior

An assessment of the effect of slow crack growth sensitivity of $MgAl_2O_4$ can be carried out using the strength values measured as a function of the loading rate (see also section 2.6.5). Graphical representations of data for the fine- and coarse-grained material are given in Fig. 57 and Fig. 58, respectively. The data were mathematically described using Eq. 25.

The derived SCG parameters for the fine-grained transparent MgAl₂O₄ are $n \sim 50$ and $D \sim 140$ MPa. For the coarse-grained spinel $n \sim 23$ and $D \sim 71$ MPa (Fig. 58) are obtained.

RT values of $n \sim 8$ to 22 have been reported for soda lime glass and $n \sim 37-50$ for Al₂O₃ [¹⁵²]. Contrary to the SCG-sensitive soda lime glass, the high *n* value of especially the fine-grained MgAl₂O₄ spinel, which is similar to that of Al₂O₃, indicates little influence of SCG at room temperature on the crack growth under standard environmental laboratory conditions.

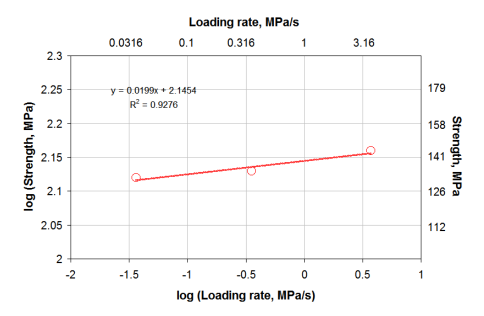


Fig. 57: *Fracture strength as a function of the loading rate for the fine-grained spinel (the standard deviation for all data points is 4%)*

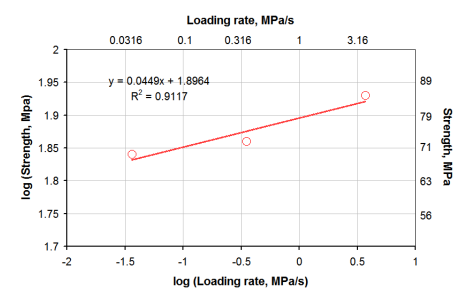


Fig. 58: *Fracture strength as a function of the loading rate for the coarse-grained spinel (the standard deviations for all data points is 3%)*

4.9.3 Strength-probability-time behavior

Based on the SPT diagram, the design stress for an acceptable failure probability can be estimated. The plots are constructed using strength, Weibull modulus and SCG parameters, as outlined in sections 2.6.4 and 2.6.5. As an example, Fig. 59 reveals that the maximum stress for a lifetime of 40 years should not exceed 56 MPa in order to warrant a failure probability of 1% for the fine-grained spinel. Based on statistical evaluation of the coarse-grained spinel data, the respective SPT diagram (Fig. 60) yields a lower maximum stress of 25 MPa for a lifetime of 40 years, warranting a failure probability of 1%.

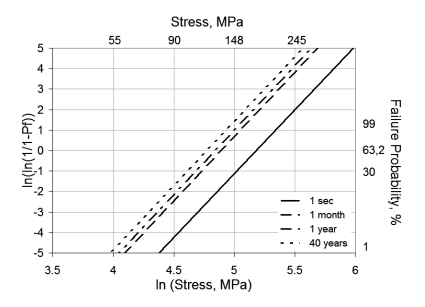


Fig. 59: *Strength–probability–time plot for the fine-grained MgAl*₂O₄ *spinel*

The SCG parameters *n* and fracture stresses determined from the SPT plot for a lifetime of 1000 h are compared in Table 11 with literature data [153] for other MgAl₂O₄ spinels.

The SPT calculations were made for the ring-on-ring specimens, according to Fig. 12. It should be noted that strength further decreases for larger component sizes, which have to be considered as outlined in section 4.9.1.

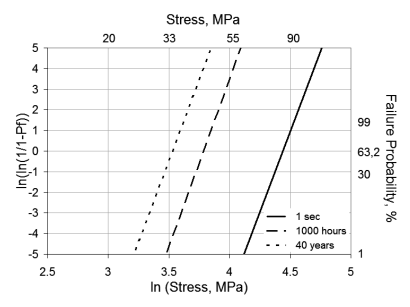


Fig. 60: *Strength–probability–time plot for the coarse-grained MgAl*₂O₄ *spinel*

Table 11. SCG parameter of spinels with different grain sizes from the present study (grey) and the literature (white) $[^{153}]$.

Grain size, µm	300	110	60	25	5
SCG, n	21	28	23	39	51
$\sigma_{ heta},$ MPa (for 1000h)*	31	35	44	52	106

* characteristic strength σ_0 for 1000h was calculated based on an SPT diagram.

The graphical presentation of these data in Fig. 61 clearly verifies that strength and slow crack growth parameters consistently decrease with increasing grain size. In fact, the grain size effect appears to be stronger for characteristic strength than for n. However, other production-related effects (different defect size distribution) might also influence the results.

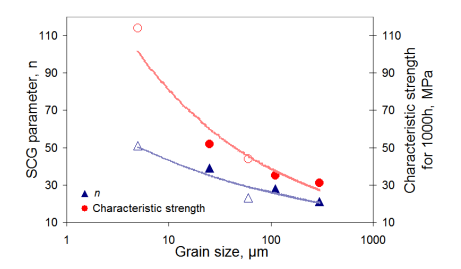


Fig. 61: Grain size dependency of SCG parameter and characteristic strength for 1000 h lifetime. Filled and unfilled circles are literature values and statistically calculated data from the present work, respectively.

5 CONCLUSION

A combination of different mechanical testing methods has been used to characterize fine- and coarse-grained spinel materials. All obtained characteristics are summarized in Table 12.

The X-ray phase analysis revealed no differences in phase composition between both spinels, displaying face centered cubic crystalline structure with lattice parameter a = 0.808 nm. Average grain sizes of ~5 µm and ~60 µm were obtained for the fine- and coarse-grained spinel, respectively, on the basis of light and electron microscopic investigations. However, X-ray and EDX analysis confirmed that even though the grain size differs and coarse-grained agglomerates (~200 – 2000 µm) exist in the fine-grained spinel, the lattice parameter and phase composition were identical, making a direct comparison of both spinels possible.

The macro-indentation testing (which was carried out for a load range from 9.8 to 98 N), revealed a hardness decrease from 17 to 13 GPa for the fine-grained spinel, whereas hardness decreases even stronger for the coarse-grained material – from 16 to 9 GPa. However, micro-indentation yielded almost identical hardness values of 16 ± 2 GPa for both spinels, in a load ranging from 0.5 to 3 N. Nevertheless, based on analysis presented in literature, it can be suggested that the coarse-grained spinel is less prone to impact damage due to the stronger indentation size effect (ISE). A detailed correlation of the measured hardness with the local microstructure of the specimens revealed that the coarse-grained material has hardness values ranging from ~12 to 17 GPa for the entire range of existing grain sizes, a data range that seems to be more narrow than for the fine-grained spinel, and, furthermore, exhibits lower average hardness values.

The fracture toughness for both spinels has been measured in a comparative approach by four different methods in order to obtain an accurate representation of the materials' behavior and to select the most appropriate and convenient method. The indentation fracture toughness (K_{IND}) appears to be less reliable than the other methods

used, since specific materials' conditions (the minimum load necessary, strong local differences in grain structure and chipping at high loads) complicate the evaluation. Nevertheless, among the relationships used, the equations suggested by Lankford et al. and Niihara et al. appear to be the most accurate, which reveal a value of $K_{IND} = 2.0 \pm 0.1$ and 1.9 ± 0.1 MPa·m^{0.5} for the fine-grained spinel, respectively. The values obtained by the other methods ($K_{ISM} = 1.9 \pm 0.2$ MPa·m^{0.5}, $K_{SENB} = 1.8 \pm 0.2$ MPa·m^{0.5} and K_{IC} (*Griffith*) = 1.6 ± 0.2 MPa·m^{0.5}) correspond well with the indentation fracture toughness and data presented in the literature. For the coarse-grained spinel, only K_{IND} and K_{SENB} values have been evaluated. In contrast to fine-grained spinel, only the indentation fracture toughness based on the Niihara equation $K_{IND} = 1.2 \pm 0.05$ MPa·m^{0.5} corresponds with the value obtained by the single-edge notch bending test $K_{SENB} = 1 \pm 0.4$ MPa·m^{0.5}. In conclusion, for both spinels the indentation fracture toughness (K_{IND}) based on the Niihara equation fracture toughness (K_{IND}) based on the Niihara equation fracture toughness (K_{IND}) based on the Niihara equation fracture toughness (K_{IND}) based on the Niihara equation fracture toughness (K_{IND}) based on the Niihara equation fracture toughness (K_{IND}) based on the Niihara equation fracture toughness (K_{IND}) based on the Niihara equation fracture toughness (K_{IND}) based on the Niihara equation ($K_{IND} = 1.2 \pm 0.05$) based on the Niihara equation ($K_{IND} = 1.2 \pm 0.05$) based on the Niihara equation ($K_{IND} = 1.2 \pm 0.05$) based on the Niihara equation ($K_{IND} = 1.2 \pm 0.05$) based on the Niihara equation ($K_{IND} = 0.5$) based on the Niihara equation ($K_{IND} = 0.5$) based on the Niihara equation ($K_{IND} = 0.5$) based on the Niihara equation ($K_{IND} = 0.5$) based on the Niihara equation ($K_{IND} = 0.5$) based on the Niihara equation ($K_{IND} = 0.5$) based on the N

Young's modulus was also determined by several methods: indentation, bending tests and impulse-excitation (IE). Impulse-excitation tests resulted in identical values at room temperature for both materials of $E_{IE} = 270 \pm 5$ GPa. At elevated temperatures (> 1000 °C) the value decreases by ~10%. Despite careful compliance correction, the bending tests yielded ~5% and ~20% lower values for the fine- and coarse-grained spinel, respectively. The lowest values were obtained by indentation testing (~25% lower). Overall, the IE and bending methods were recognized as most reliable and the data corresponded well with literature [¹,⁵⁸⁻⁶¹].

Surface quality, effective area, inherent defects and structure of the material strongly influence the fracture strength. In the ring-on-ring bending test, characteristic strengths of $\sigma_0 = 169 \pm 3$ MPa and $\sigma_0 = 77 \pm 1$ MPa were obtained for fine- and coarse-grained spinel, respectively. Subsequent fractographic analysis showed mixed trans- and intercrystalline fracture in fine- and pure intergranular fracture in the coarse-grained spinel. Due to the presence of coarse-grained agglomerates in the fine-grained spinel, crack growth is affected by these inhomogeneities. However, there is no correlation of fracture stress with size and volume content of the agglomerates. Furthermore, Hertzian indentation has proven to be a non-destructive characterization approach, capable of estimating the local strength.

The experimentally obtained mechanical parameters were statistically analyzed using a Weibull distribution, a subcritical crack growth approach and strength-probability-time prediction for long-term reliability assessment. In fact, the obtained characteristic strength σ_0 and slow crack growth exponent *n* are almost a factor of 2.5 times higher for fine-grained spinel, whereas the Weibull modulus is a factor of 2 lower, compared to the coarse-grained spinel.

Overall, it can be concluded that the fine-grained spinel has a moderate strength but a lower sensitivy to slow crack growth, whereas the strength is less sensitive to the specimen size for the coarse-grained spinel, is less prone to impact damage, and has a higher Weibull modulus.

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Table

		Strength, MPa	ИРа		в И) СРа		Toughness MPa•√m	mess √m		Young'	Young's Modulus, GPa	, GPa
	Average	Weibull parameter	ull eter	SCG	,eeenb ,eeepb ,eeepb ,eeepb ,eeepb ,eeepb ,eeepb ,eeepb ,eeepb ,eeepb ,eeepb ,eeepb ,eeepb ,eeepb ,eeepb ,eeepb ,eeepb ,eeebb ,eebbb ,eebbb ,eebbb ,eebbb ,eebbb ,eebbb ,eebbb ,eebbb ,eebbbb ,eebbbb ,eebbbb ,eebbbb ,eebbbb ,eebbbbb ,eebbbbb ,eebbbbbbb ,eebbbbbbbb	K_{IND}	K	K	K_{IC}	$E_{biax.}$	E_{IT}	E_{IE}
	σ_f	σ_{0}	ш	и		(Niihara)	WSIX	A SENB	(Griffith)	GPa	GPa	GPa
Fine-grained $(\sim 5 \ \mu m \ g.s.)$	155±38	169±3	5±2	51	51 16±2	1.9±0.1	1.9±0.2	1.8±0.2	1.8±0.2 1.6±0.2	260±5	210±10	270±5
Coarse-grained $(\sim 60 \ \mu m \ g.s.)$	73±9	77±1	10±2	23	16±2	77±1 10±2 23 16±2 1.2±0.05		1±0.4		242±13	225±10	270±5

CONCLUSION

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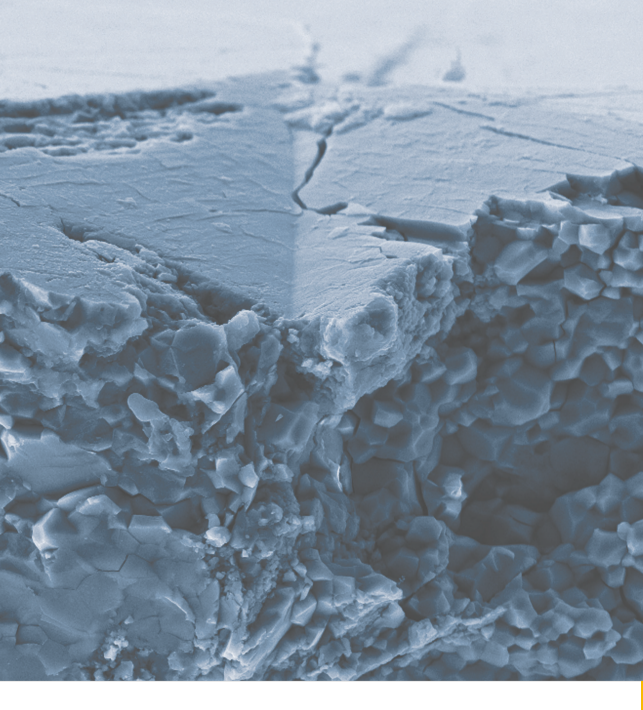
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