Mechanism and Kinetics of Oxidation of ZrN Ceramics

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Abstract
Oxidation of ZrN ceramics from 973-1373 K under static conditions reveals parabolic rate behaviour, indicative of a diffusion-controlled process. In-situ high temperature XRD found the oxidation mechanism begins with destabilisation of ZrN through formation of a ZrN$_{1-x}$ phase with oxide peaks initially detected at around 773 K. The zirconium oxide layer was found to be monoclinic by in-situ XRD with no evidence of tetragonal or cubic polymorphs present to 1023 K. Samples oxidised at 1173 and 1273 K underwent slower oxidation than those oxidised at 973 and 1073 K. This change in oxidation rate and hence mechanism was due to formation of a denser c-ZrO$_2$ polymorph stabilised by nitrogen defects. This N-doped dense ZrO$_2$ layer acts as a diffusion barrier to oxygen diffusion. However, at an oxidation temperature of 1373 K this layer is no longer protective due to increased diffusion through it resulting in grain boundary oxidation.

Keywords: Reaction Rate, Oxynitride, Nitride, Oxidation, Zirconium

1. Introduction
Zirconium nitride is being considered for use in advanced nuclear power plants as inert matrix fuel (IMF) or as accident tolerant fuel particle coatings due to its high thermal conductivity (45-50 Wm$^{-1}$K$^{-1}$ [1, 2]), low neutron capture cross section and chemical compatibility with existing fuel cycle technology [3]. Due to this increased interest in ZrN, recent work has focused on assessing thermophysical properties of ZrN and mixed phases of actinide nitrides dispersed in ZrN [1, 2, 4–10]. However, there have been few studies on the oxidation behavior of ZrN which is an important factor to be considered especially under accident conditions such as occurs in a loss of coolant accident (LOCA).

The majority of oxidation studies of ZrN have been performed on thin films due to the extensive use of ZrN as a hard coating on cutting tools [11–13]. Krusin-Elbaum and Wittmer [13] studied the oxidation kinetics of ZrN thin films (60 nm -1 µm) reporting the activation energy of oxidation to be 241 ± 10 kJ mol$^{-1}$ from 748-923 K with the oxide layer comprising monoclinic and cubic ZrO$_2$. Panjan et al.[12] showed the oxidation of ZrN coatings with thicknesses of 300 µm to have an activation energy of 229 kJ mol$^{-1}$ from 773-1123 K and obey a parabolic rate law, in agreement with Krusin-Elbaum and Wittmer.[13] Both studies show a plateau region after initial oxidation with both authors reporting parabolic rate behavior and it was suggested the diffusion of oxygen through the oxide layer is the rate limiting step. Caillet et al. [14] studied the oxidation of 30 µm ZrN coatings from 823-973 K observing linear kinetics suggesting oxygen diffusion through the oxide layer does not affect the rate, however a significant mass loss was
observed around 823 K. The authors propose nitrogen is initially lost leaving hcp-Zr at the surface prior to ZrO$_2$.

Zirconium oxynitrides have been produced previously by mixing of ZrO$_2$ and ZrN powders \[15–17\] or by nitriding ZrO$_2$ under NH$_3$ gas.\[18\] Three oxynitride polymorphs have been reported by Gilles et al. \[17\] Zr$_2$ON$_2$, Zr$_7$O$_8$N$_4$ and Zr$_7$O$_{11}$N$_2$ which are designated $\gamma$, $\beta$ and $\beta'$ respectively. It can be envisaged that the formation of an oxynitride phase would be formed at the oxidation interface of ZrN, similar to behaviour observed on oxidation of Si$_3$N$_4$.\[19, 20\]

ZrC has similar chemical properties and crystal structure to ZrN and its oxidation kinetics and mechanism in the bulk are better understood \[21–24\] with onset of oxidation occurring between 653-750 K\[25\] similar to that of ZrN.\[13\] Oxidation of ZrC above 743 K occurs via the following sequence of steps: \[21\]

- Formation of a ZrO$_x$C$_y$ phase surface, which proceeds to form amorphous ZrO$_2$ and free C.
- Crystallites of cubic zirconia (c-ZrO$_2$) nucleate and grow to form a dense oxide layer with free carbon, which stabilises the c-ZrO$_2$.
- Oxygen then diffuses through this oxide layer to the free carbon producing CO$_2$ which escapes via any existing cracks or pores leaving behind voids and pores in the zirconia layer which, with less carbon left to stabilise transforms to monoclinic zirconia (m-ZrO$_2$) with small amounts of tetragonal polymorph (t-ZrO$_2$). \[25\]

ZrN was expected to show similar oxidation properties as ZrC and the aims of this work were to examine the mechanism of oxidation of bulk ZrN samples, determining phases present in the oxidation layer and investigate the existence of different layers at the reaction interface, such as oxynitride or high temperature ZrO$_2$ polymorphs as observed with ZrC ceramics.\[25\] Oxidation temperatures were chosen as to be able to examine the initial to intermediate stages of oxidation.

The higher temperatures (1073-1373 K) are also in the range of the expected nominal operating temperature and initial temperature increases in the core following a LOCA.\[26\] Static atmosphere was used to aid examination of the initial and intermediate steps of oxidation and also as this is envisaged as being more akin to a LOCA whereby the coolant system fails but the nuclear reactor core remains intact.

2. Experimental

Commercially-available ZrN powder (Sigma Aldrich, 1-2 μm, $\geq$ 99.0%) was densified using spark plasma sintering (HP D/25/1, FCT systeme GmbH, Rauenstein, Germany) in a graphite die (30mm diameter), under vacuum (0.5 mbar). Sintering was carried out for 10 min at 2373 K (+100 K/min) under 50 MPa uniaxial pressure. The resulting bulk ceramic samples had a density of 7.13 ± 0.01 g/cm$^3$ calculated from the average of three measurements performed by the Archimedes method and a theoretical density of 97.7 ± 0.1%.

In-situ phase analysis of ZrN powder (Sigma Aldrich, 1-2 μm, $\geq$ 99.0%) under static air was done using high temperature (up to 1023 K) X-ray diffraction (HT-XRD) (X’pert MD, Phillips, Amsterdam, Netherlands) in a furnace using a Pt heating strip (HDK 2.4, Buhler, Tubingen, Germany) at 40 kV and 40 mA using Ni-filtered CuK$\alpha$ radiation between room temperature and 1023 K, with scans performed every 12 K between 673-1023 K. Due to the nature of the in-situ XRD requiring thin layers of sample experiments were limited to powders.

Ceramic samples with dimensions 3×3×3 mm hand abraded with 600 grit SiC paper were oxidised under static air using thermogravimetric analysis coupled differential thermal analysis
(TGA/DTA) to measure the change in weight as a function of time (STA-449-F1, Netzsch GmbH, Selb, Germany) at 5 temperatures; 973, 1073 and 1173, 1273 and 1373 K (+20 K/min) for a dwell time of 5 h. Change in mass was continually recorded using the micro-balance of the TGA. Oxidation using the TGA ensures more reliable results in weight change as any oxide scale spallation remains in the crucible and there is no excess weight gain from the cooling of the furnace at the desired time points. Experiments were performed once at each isothermal oxidation temperature so results are directly comparable. Plots of $\frac{\Delta W}{A}$, $(\frac{\Delta W}{A})^2$ and $(\frac{\Delta W}{A})^3$ against time were made to determine which equation gave the the best linear fit according to equations 1, 2 and 3 (where $\Delta W$ is the change in weight (mg), $A$ is the area of the pellet (cm$^2$), $k_l$, $k_p$ and $k_c$ are the linear, parabolic and cubic rate constants respectively and $t$ is the time (s)). Errors of $\frac{\Delta W}{A}$ are given as the percentage error of the TGA/DTA instrument compounded with the error from the area measurements of the pellets.

$$\frac{\Delta W}{A} = k_l t$$ (1)

$$(\frac{\Delta W}{A})^2 = k_p t$$ (2)

$$(\frac{\Delta W}{A})^3 = k_c t$$ (3)

Microstructural characterisation of the oxidised, polished pellets was performed using secondary electron imaging (SEI) in an SEM (JSM-6400, JEOL, Tokyo, Japan) equipped with energy dispersive spectroscopy (EDS) detector for chemical analysis (ultra thin polymer window, INCA, Oxford instruments, Oxford, UK). Focused ion beam (FEI, Helios Nanolab 600, OR, USA) was used to prepare samples and transmission electron microscope (TEM) analysis was performed on the oxidised ceramics (HRTEM FX2000, JEOL, Tokyo, Japan). Selected area diffraction (SAD) patterns were indexed by matching $d_{hkl}$ values with reference patterns and calculated SAD patterns using SingleCrystal software (version 2.2.9). Lattice parameters were calculated from the indexed SAD patterns along with the measured $d_{hkl}$ spacing’s using equation 4 and contain a 2% error due to the small (0.5-5$^\circ$) angle of diffraction.[29]

$$a = d \sqrt{h^2 + k^2 + l^2}$$ (4)

3. Results and discussion

3.1. Oxidation kinetics

In-situ XRD of ZrN powder during oxidation is presented in figure 1. Monoclinic ZrO$_2$ is initially detected at 773 K after no dwell time and was the only polymorph detected during the oxidation process which as expected since the tetragonal and cubic forms are both high temperature ($\geq 1450$ K) polymorphs. However, this is not in agreement with observations of a very low intensity reflections identified as a minor cubic ZrO$_2$ polymorph detected by glancing angle XRD of oxidised thin films by Krusin-Elbaum and Wittmer [13]. It should be noted that the observation of low amounts of cubic ZrO$_2$ would be difficult by in-situ XRD due to the differences in experimental technique with glancing angle diffractometers. Peak intensity is also lower in in-situ XRD compared to typical diffractometers due to the scattering from the chamber window of the furnace between the source and detector. Figure 2 shows an excerpt of the (200) ZrN reflection which reveals that initially the peak shifts to lower 2$\theta$ angles, corresponding to
an increase in lattice parameter due to thermal expansion until around 673 K. Above 673 K
the peaks begin to shift to higher 2θ angles suggesting a decrease in lattice parameter which
could arise from loss of nitrogen from the lattice resulting in a ZrN$_{1-x}$. Further increase of
temperature leads to the peak splitting, resulting in a peak consisting of two main components
(observed for all ZrN reflections). This suggests the existence of two or more ZrN phases, such
as a substoichiometric ZrN$_{1-x}$ phase and Zr-O-N phase. Incorporation of oxygen into the ZrN
lattice to form an oxy nitride with the same crystal structure would result in the formation of a
smaller lattice resulting in the peak shift to higher 2θ. This agrees with the observations of Gendre
et al.[30] who showed the lattice size of ZrC decreased with oxygen content to a stoichiometry of
around ZrC$_{0.79}$O$_{0.13}$.

The change in weight normalized for the surface area of the ceramic samples as a function
of time is shown in figure 3 which reveals a non-linear relationship (equation 1) for all samples
(indicated by R$^2$ values $\leq 0.99$ for a linear fit in figure 3), except for the sample oxidised at 1273
K which fits well to a linear relationship. However, this may be because the oxidation rate is
still in the linear region and longer oxidation times may show parabolic rate behaviour. Figure 4
shows the square of the weight gain normalised for surface area as a function of time for each
isothermal oxidation temperature with all R$^2$ values $\geq 0.99$ showing a good fit. All samples had
a R$^2 \leq 0.97$ for a cubic relationship (equation 3) showing a poor fit. A parabolic rate is expected
for the samples due to increased diffusion distance for oxygen through the growing oxide layer
to the ZrO$_2$/ZrN reaction interface. It can be seen from figure 3 and plotting the rate constant
for each temperature (figure 5) that the samples oxidised for 5 h at lower temperatures (973 and
1073 K) showed more weight gain than those oxidised for 5 h at 1173 and 1273 K. The rate
constants in figure 5 show a rapid increase between 973 and 1073 K followed by a large decrease
at 1173 K and then no significant difference between samples oxidised at 1173 and 1273 K. At
the higher temperature of 1373 K however, the rate of oxidation is the fastest observed for all
isothermal oxidation temperatures. The proposed change in mechanism between low and high
temperature oxidation will be discussed further in section 3.3.

3.2. Oxidation layer characterisation

The ceramic sample oxidised for 5 h at 973 K (figure 6) an outer layer of brittle, porous
ZrO$_2$ was formed which spalled away from the surface of the sample, revealed by loose powder
present when removing the sample from the crucible. Figure 6a also shows light regions with
increased porosity further into the bulk from the surface that EDS revealed contained oxygen
and nitrogen, indicating the oxidation layer penetrated further than the surface layer. This is
more clearly observed in the sample oxidised at 1073 K for 5 h (figure 7) where a dense but
cracked layer of ZrO$_2$ at the surface of the sample is observed on top of a cracked layer with
mixed porosity which EDS also showed contained oxygen. SEIs of the samples heated to 1173
and 1273 K for 5 h are shown in figures 8 and 9 respectively. Figure 8 reveals three layers, EDS
of the first two revealed only Zr and O, while the third was unreacted bulk ZrN. The presence
of carbon arises from the epoxy resin used to mount the samples. The SEIs in figure 9 reveal
that the grains closer to the surface (nearer to the oxide layer) are readily visible however, grains
towards the bulk are not. Line scans of figure 9b reveal the Zr counts decrease in the crack in
the outer oxide layer, however, do not decrease significantly at the revealed grain boundary. This
suggests the grains have been revealed by oxygen diffusion along the grain boundaries nearer to
the surface. A decrease in oxide layer thickness (around 20µm thinner) between the samples
oxidised at 1073 and 1173 K for 5 h was observed. The oxide scale thickness did not show a
significant increase for the samples oxidised at 1173 and 1273 K for 5 h. SEIs (figure 10) reveal
the sample oxidised at 1373 K for 5 h showed extensive grain boundary oxidation in the bulk.
The oxide scale is also cracked and spalled away from the sample on handling, similar to the sample oxidised at 973 K for 5 h.

TEM of a FIB section of the intermediate layer observed in the sample oxidised at 1173 K is shown in figure 11a. The dark phase is a thick region of monoclinic ZrO$_2$ as characterised by XRD (figure 1) and corresponds to the surface of the oxidised sample. The bulk of the sample is ZrN and the SAD pattern from this region matched ZrN (figure 11b which indexed as the [001]ZrN zone axis). However, the calculated lattice parameter of 4.400±0.090 Å is smaller than the literature value given at 4.574 Å.[31] The decrease in lattice size may be due to nitrogen vacancies and oxygen content causing the lattice to shrink. SAD patterns of the intermediate grain between the bulk ZrN grain and the surface ZrO$_2$ are shown in figures 11c and d, which has a width of around 1 µm and agrees well with the width of the layer in figure 8. The SAD pattern in figure 11d indexed as the c-ZrO$_2$ [103] zone axis as the ratio of the (311)/(002) d-spacings (1.678) matches with the cubic pattern (1.658) as opposed to the ratio of 1.728 in the tetragonal ZrO$_2$ [103] zone axis. The lattice parameters calculated from the SAD patterns in figures 11c and d were 4.877±0.100 and 4.877±0.100 Å respectively, which although ≈ 0.2 Å smaller than the cubic ZrO$_2$ lattice parameter of 5.090 Å [32] match better than with the ZrN lattice parameter of 4.574 Å [31]. Figure 12a shows a TEM image of a FIB section a grain boundary from the sample oxidised at 1373 K for 5 h, which reveals the material between the ZrN grains in figure 10 to consist of sub-micron grains, intergranular cracking can also be seen. The SAD pattern of the sub-micron grains shown in figure 12 indexed as the cubic ZrO$_2$ [111] zone axis and had a lattice parameter of 4.850±0.100 Å. Although nitrogen has a larger atomic radii than oxygen due to its weaker effective pull on its outer electrons a decrease in observed cubic ZrO$_2$ lattice parameter may be due to oxygen vacancies formed by substitution of oxygen with nitrogen. Evidence for ordering of oxygen vacancies and nitrogen impurities can also be observed from the forbidden reflections present in figures 11c,d and figure 12 which have d-spacing values which are half of the fundamental reflection spots, suggesting they are from superlattice reflections and not from neighbouring crystals of differing lattice size. Similar forbidden reflections and stabilisation of c-ZrO$_2$ by the addition of ZrN were observed by Van Tendeloo and Thomas [15]. The authors state that the vacancies introduced by replacing oxygen with nitrogen order along one of the three fold axes which lowers the symmetry of a rhombohedral Zr$_7$O$_{11}$N$_2$ phase which is closely related to the cubic fluorite structure allowing the forbidden reflections. It is plausible a similar phase may be observed in this work, however, only the observation of stabilisation of c-ZrO$_2$ was made with certainty.

3.3. Discussion of Oxidation Mechanism

Parabolic rate behavior was observed for all oxidation temperatures except the sample oxidised at 1273 K for 5 h, indicative of a diffusion controlled rate limiting step. The observed decrease in rate of oxidation between 1073 and 1173 K along with the difference in morphology of the oxidation layer cross sections (e.g. figure 7 compared to figure 8) and the presence of a high temperature c-ZrO$_2$ phase at the oxidation layer (figures 8, 11 and 11) suggest the mechanism of oxidation changes with temperature and a schematic is shown in figure 13. At temperatures up to 1073 K a porous oxide layer forms, which would readily allow oxygen diffusion through pores and voids and nitrogen (or NO$_x$ species) diffusion out of the sample. The increase in oxidation rate (figure 5) is consistent with the porous microstructure of the oxide layers (figures 6 and 7). At oxidation temperatures of 1173 K and 1273 K a dense intermediate oxide layer forms which inhibits the diffusion of oxygen and therefore the rate of oxidation. This dense layer is revealed to be c-ZrO$_2$ by TEM results of the intermediate layer (figure 11). As the sample cools the oxide layer transforms to the monoclinic polymorph creating the cracks present in the dense ZrO$_2$ polymorph at the surface (figures 8 and 9) due to the volume change associated
with the polymorphic transformation. However, the sample oxidised at 1373 K for 5 h showed a rapid increase in rate of oxidation. It can be seen from the SEIs (figure 10) that there has been extensive grain boundary oxidation which would result from the increased diffusion of oxygen through the oxide scale.

Shimada and Ishii [21] describe a change in mechanism of ZrC oxidation with increasing temperature. At higher temperatures (≥743 K), submicron c-ZrO$_2$ crystals nucleate from the amorphous ZrO$_2$ + C layer and this forms a compact dense layer whose formation becomes the rate determining step for oxygen diffusion. However, at higher temperatures (823 K) and longer oxidation times the small crystallites grow and the volume expansion causes intergranular fracture providing paths for oxygen diffusion in and CO$_2$ diffusion out. Molar volumes of ZrN, m-ZrO$_2$ and c-ZrO$_2$ are presented in table 1 along with the thermal expansion data for ZrN and yttria-stabilised ZrO$_2$. Thermal expansion data for yttria stabilised ZrO$_2$ is assumed to be similar to the stabilised ZrO$_2$ polymorph in this work. From the values in table 1 it can be seen the associated volume change between phases and the thermal expansion mismatch between stabilised ZrO$_2$ could both provide sources of stresses and result in intergranular fracture. Puclin and Kaczmarek [33] examined the growth in crystal size of ZrO$_2$ reporting a rapid increase in size of t-ZrO$_2$ at 1223 K and m-ZrO$_2$ at 1273 K which can also be envisaged as a source of stress and cracking due to the growth of submicron size of the c-ZrO$_2$ grains in the grain boundary.

From the observation of a thin (≤1µm) c-ZrO$_2$ layer in the ZrN samples in this work (figure 8) and small grains (≤1µm) observed by TEM (figure 11) a similar rate limiting step can be envisaged for ZrN, which causes a decreases in the rate of oxidation between the samples oxidised at 1073 and 1173 K. However, these crystallites then grow and could transform to m-ZrO$_2$ causing cracking between the grains, as observed in 12a, where it can be seen the crack follows what was the ZrN grain boundary in the oxide layer and slightly into the bulk. Berkowitz-Mattuck [34] also observed that oxidation of ZrC samples at temperature ≤1470 K resulted in porous m-ZrO$_2$ which fractured due to intergranular oxidation, however above 1470 K a denser ZrO$_2$ layer formed slowing diffusion of oxygen to the bulk attributed to sintering effects on the oxide layer. The observations of denser outer layers of ZrO$_2$ of the samples oxidised 1173 and 1273 K (figures 8 and 9) agree well with the dense ZrO$_2$ layer reported by Shimada and Ishii [21] and Berkowitz-Mattuck. [34] Simultaneously the formation of dense scales of ZrO$_2$ have been proposed to provide effective oxidation resistance in ZrB$_2$-SiC [35] and ZrC-SiC [36] composites.

Different mechanisms are thus proposed for the low temperature (≤1073 K) and higher temperature (≥1173 K) oxidation of ZrN. The oxidation mechanism ≤1073 K can be described in several steps, initially oxygen diffusion through the surface ZrN grain boundaries results in a ZrO$_2$N$_y$ phase which then forms m-ZrO$_2$. The formation of this oxide layer results in a diffusion-controlled process as observed by the parabolic kinetics shown in figure 4. Stresses from the m-ZrO$_2$ grains between the ZrN grains result in intergranular fracture and the creation of a porous oxide layer as observed in figure 6, similar to the observation of Berkowitz-Mattuck for ZrC.[34] Figure 7 also reveals lighter regions in the bulk of the ZrN which correspond to oxygen-rich areas formed from grain boundary diffusion of oxygen that have not yet caused fracture. The creation of these grain boundary fractures opens channels for further ingress of oxygen and removal of nitrogen advancing the rate of oxidation, this is shown schematically in figure 13 labelled ≤1073 K.

Oxidation of ZrN at higher temperatures, however, proceeds by the formation of a high temperature c-ZrO$_2$ polymorph comprised of submicron grains. This high temperature phase is stabilised by the substitution of oxygen with nitrogen forming oxygen vacancies. As the oxidation of ZrN is exothermic [37], it is possible the local temperature at the interface will be higher than the oxidation temperature. This increase in temperature may give rise to sintering of the c-ZrO$_2$ crystals similar to effects observed in the oxidation of ZrC by Berkowitz-Mattuck [34] however,
the nominal sintering temperature of ZrO$_2$ is around 1673 K. This dense c-ZrO$_2$ is also similar to that reported by Shimada and Ishii [21]. The dense oxide layer inhibits oxygen’s ingress and therefore reduces the rate of oxidation, providing a diffusion barrier resulting in parabolic behavior and a lower oxidation as shown by the rate constants in figure 5. However, from figure 9, grains are more readily observed suggesting greater potential for oxygen diffusion via grain boundaries. Expansion of the c-ZrO$_2$ crystallites or transformation to monoclinic as nitrogen defects are removed results in the cracks present in the oxide layers in figures 8 and 9. The sample oxidised at 1373 K for 5 h showed an increase in oxidation rate and this is attributed to failure of the dense oxide scale as, although oxygen diffusion through the dense zirconia layer will be slow, it will increase with temperature. This is observed as the increase in grain boundary oxidation of samples oxidised at 1273 and 1373 K for 5 h (figures 9 and 10 respectively). Thus, eventually the protective oxide scales in the samples oxidised at 1173 and 1273 K will fail with time as oxygen diffuses further through the dense oxide layer. Intergranular cracking observed in the TEM image of the sample oxidised at 1373 k for 5 h (figure 12) will create channels for further oxygen diffusion into the sample.

4. Conclusions

The oxidation kinetics and mechanism of bulk ZrN ceramics between 973 and 1373 K have been studied. In-situ powder XRD reveals oxidation begins by removal of nitrogen from the ZrN lattice (forming two or more phases with smaller lattice size indicating either substoichiometric ZrN or a ZrN phase with oxygen defects) and proceeds by formation of a porous m-ZrO$_2$ phase at temperatures up to 1023 K. Oxidation rates decreased at temperatures at or above 1173 K attributed to formation of a higher energy c-ZrO$_2$ phase which is proposed to slow oxygen diffusion by forming a dense layer, limiting the diffusion of oxygen to the bulk. At 1373 K however, this barrier to oxygen diffusion fails due to increased oxidation of grain boundaries and intergranular cracking creating more channels for oxygen diffusion. The nature of the protective scale observed for samples oxidised at 1173 and 1273 K for 5 h needs further investigation, examining longer oxidation times and oxygen pressures.

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References


1. In-situ XRD characterisation of ZrN powder from RT-1023K with scans performed every 100 K between RT-673 K and every 12 K between 673-1023K with no dwell time at 1023 K. Peaks indexed with monoclinic ZrO$_2$ [40] and ZrN [31] ...

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<th>Phase</th>
<th>Molar volume ($V_m$) (cm$^3$ mol$^{-1}$)</th>
<th>Coefficient of thermal expansion at 1273 K ($10^{-6}$ K$^{-1}$)</th>
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**Table 1**: Molar volumes and thermal expansion coefficients of ZrN, 3 mol% yttria-stabilised ZrO$_2$, and m-ZrO$_2$ [40–42]