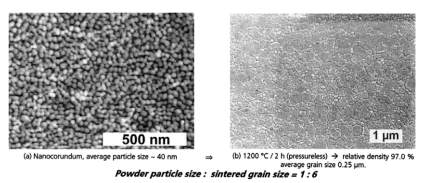
# TP3 - Ceramics: Sintering and Microstructure

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#### 1. Introduction

After the initial molding of the ceramic, whether by slip casting or dry pressing, it is still necessary to densify the compacted powder samples (green bodies) to form a continuous 3D structure and thus to get ceramic pieces appropriate for the selected application. Usually this is done by a sintering process which is a consolidation and densification step of the granular compact through the action of heat. During this stage, the microstructure of the samples evolves (see Figure 1).



**Figure 1** *Sintered Powder and microstructure. Source Krell et al.*[8].

Pores, second phases and grain size have all an important influence on many of the final properties of ceramics. In general, the porosity has a more pronounced effect than a second phase or the grain size, because most of the desired properties (e.g. elasticity or thermal conductivity) are zero or close to zero in the pores. Thus porosity is often detrimental to important properties such as elastic modulus, tensile strength, hardness or the thermal and electrical conductivities. The desired final properties depend on the final application of the sintered piece. For example, for a hip replacement a very high mechanical strength is required, so the residual porosity of the sintered part should be very low. However, for a catalyst, a high specific surface area is usually essential and therefore the porosity after sintering should be high while still ensuring a good cohesion of the piece.

There are three types of sintering with different densification mechanisms:

- 1. Solid phase sintering: all components remain solid throughout the sintering. The densification is carried out by a change in shape of the grains. Mass transport occurs by volume and grain boundary diffusion.
- 2. Liquid phase sintering: formation of a viscous liquid (usually a eutectic with a low melting point) that fills the pore spaces of the initial green body (e.g. porcelain). Densification occurs mainly by dissolution and re-precipitation of the solid which allows a rapid mass transport.
- 3. Reactive sintering: two or more constituents react during sintering. The densification is carried out by the formation of a new compound.

From a technical perspective, the densification during sintering can be improved by applying an external force. The most common techniques are the application of a uni-axial pressure (HP = hot pressing) or isostatic pressing (HIP = hot isostatic pressing). Additionally, there is a variety of

techniques to influence the microstructural characteristics of the final piece by changing the heating and cooling rates. An example is spark plasma sintering (SPS) where very high heating and cooling rates are achieved with the help of a pulsed electric field which leads to a high density in a relatively short period of time (5-10 minutes) while keeping a relatively small grain size. [12]

The subject of this lab is about natural, solid phase sintering (i.e. heating in a conventional oven without applying an external force). Various factors influencing sintering will be discussed. The objective is to understand the driving force and the mechanisms of sintering and to get an idea of how they can be influenced. The influence of the particle size and of the addition of dopants on sintering will be studied. As the experimental methods can only provide limited information on the topics we will use digital modeling techniques to better understand what happens during sintering. To follow the progress of sintering we use a dilatometer which tracks dimensional changes in a piece during sintering. The density and residual porosity of the final piece is then studied with an experimental method called the immersion or Archimedes method.

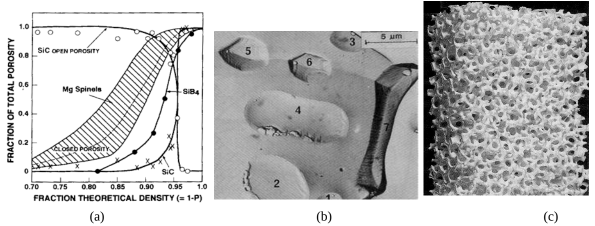
# 2. Density and Porosity Measurements

#### 2.1. Density and Porosity - Definition

As mentioned previously the porosity has a huge impact on the final properties of a sintered ceramics. However, the total volume of the porosity is insufficient to understand the sintering behavior and the final properties of the material. There are several additional characteristics that will influence the porosity:

- 1. The connectivity of the pores: there exists open and closed porosity. In the first case, the pores communicate with the outside of the material, they can theoretically be filled with a fluid by immersion of the piece. In the case of closed porosity the pores are isolated inside the material.
- 2. The pore size: the pore size is important for the final properties as well as during the sintering as small pores are much more difficult to eliminate than large ones.
- 3. The shape of pores: the pores can have different shapes which influence their behavior. Open pores are typically thin, elongated and irregularly shaped. The closed pores are typically more equiaxed.
- 4. The distribution of the porosity: the global special distribution of the porosity can be of importance. For example during the production of ceramic layers, the porosity is not always uniform throughout the thickness of the layer.

The evolution of fractions of open and closed porosity versus the fraction of theoretical density is shown in Figure 2a) for different spinels (SiC and SiB<sub>4</sub>). This shows the replacement of the open porosity by closed porosity. On a fracture surface (Figure 2b)) we can see different types of pores for  $Al_2O_3$ : flat (1-4), elongated (4 and 7) and of more complex shape (5-6). If one maximizes the porosity for a given application, one can obtain a foam (Figure 2c)). Such foams are typically used for filtration or as a catalyst because of their high surface area.



**Figure 2** Illustration of porosity: (a) evolution of open and closed porosity. (b) different shapes of pores. (c) ceramic foam with high porosity.

#### 2.2. Immersion Method / Archimedes Method

The absolute densities calculated from crystalline structures or measured using pycnometry [6] as well as the dry, wet and immersed masses can be used for the calculation of apparent densities and the volume fractions of the open and the closed porosity. The solvent used during this TP is deionized water without additive. However the same procedure can also be done with alcohol or other solvents wetting the material studied. In the case of alcohol, solvent evaporation can greatly affect the quality of measurements due to evaporation and the time between the different stages of measures must be strictly controlled.

In general one can distinguish between two different types of porosities:

- Open porosity,  $\mathbf{p_0}$ : pores that communicate with the outside of the material, they can theoretically be filled with a fluid.
- Closed porosity,  $\mathbf{p_f}$ : the pores are isolated inside the material

This means that the apparent volume  $V_{app}$  is equal to the sum of different parts of the sample: The actual volume of the solid  $V_s$ , the volume of the open porosity,  $V_{po}$  and the volume of the closed porosity  $V_{pf}$  (see equation (1)).

$$V_{app} = V_s + V_{po} + V_{pf} \qquad (1)$$

If we assume that the solvent wets the entire surface of the open porosity, the open porosity can be estimated from the impregnated  $(m_2)$  compared to the dry mass  $(m_1)$  divided by the water density  $\rho_{eau}$  (equation (2)).

$$V_{po} = \frac{m_2 - m_1}{\rho_{eau}} \tag{2}$$

The volume of the closed porosity can be determined via the buoyancy  $P_A$  or the immersed mass  $m_3$  (equation (3)): We consider that this force is exerted on the volume of the solid ( $V_S$ ) with a material dependent density  $\rho_{abs}$  (e.g. 3987 g/cm3 for alpha alumina) and on the closed pore volume ( $V_S$ ) with essentially zero density (equation (4)). This means that the volume of the closed pores can be calculated according to equation (5).

$$P_{A} = m_{3} \qquad (3)$$

$$V_{s} + V_{pf} = \frac{P_{A}}{\rho_{eau}} \qquad (4)$$

$$V_{pf} = \frac{P_{A}}{\rho_{eau}} - V_{s} = \frac{m_{3}}{\rho_{eau}} - \frac{m_{1}}{\rho_{abs}} \qquad (5)$$

The percentage of open pore volume (Po) and closed (Pf) can be calculated as follows:

$$P_o = \frac{V_{po}}{V_{app}} \cdot 100 \qquad P_f = \frac{V_{pf}}{V_{app}} \cdot 100$$

And finally the effective density of the sample  $\rho_2$  (closed pores included ....) is:

$$\rho_2 = \frac{m_1}{V_s + V_{pf}} = \frac{m_1}{P_A} = \frac{m_1 \cdot \rho_{eau}}{P_A}$$

#### Samples

The ceramic studied is alumina (chemical formula  $Al_2O_3$ ) with and without dopants. We will measure a series of three sintered pellets (natural sintering in air) at three different temperatures will be used:  $1400 \,^{\circ}\text{C}$ ,  $1500 \,^{\circ}\text{C}$  and  $1700 \,^{\circ}\text{C}$  (ramp  $10 \,^{\circ}\text{C}$  / min).

#### Operation mode

The liquid used for immersion must be stable and controlled. Water must be at a constant temperature to avoid variations in density, this is why the water used for soaking the samples (cooler than the ambient temperature) should not be used. Number the pellets using a felt pen (water resistant).

- 2. Set the balance to zero (tare) with the immersion basket in water. Weigh the dry samples (dry mass, m<sub>1</sub>)
- 3. Place the samples in a small beaker and create a void
- 4. At the same time put enough water into the beaker to immerse the samples completely
- 5. Keep the void until no more air bubbles are observed (~ 30 min)
- 6. Take the samples out of the water and dry them carefully then weigh them (impregnated mass m<sub>2</sub>). When working with alcohol, e.g. isopropanol this last step must always be done in the same time e.g. within 30 seconds.
- 7. Tare the balance and then weigh the sample in water (immersed mass, m<sub>3</sub>) (can be repeated 2 or 3 times to check reproducibility). Measure the temperature of the water for each sample.
- 8. Calculate the apparent density, the relative density and the effective, open and closed porosity according to the equations above. Use the value of the density of water from the table given in the annex.

## 3. Sintering

#### 3.1. Driving force of the sintering

The driving force for sintering is the reduction of the interfacial free energy of the system, both by replacing the solid-gas interfaces (surfaces) by grain boundaries (densification), and by reducing the

ratio of the interfacial area per volume of the grains (grain growth). Thus there are always at least two phenomena that compete during sintering: densification and grain growth.

Densification can be looked at at three different scales:

- Global scale
- Microstructure scale (grain)
- Atomic scale

On a global scale the densification is due to a minimization of the surface energy by replacing the solid-gas interfaces by grain boundaries. In other words the overall driving force for densification is the global surface energy reduction.

On the scale of the microstructure, the driving forces for mass transport are the differences in pressure and concentration gradients of point defects (e.g. vacancies) due to differences in radius of curvature in the microstructure. The relationship between pressure and the radius of curvature can be illustrated by imagining an elastic membrane that can stretch and bend under the influence of pressure at the place of the grain surfaces. A flat membrane exerts no pressure on either side (conversely, a membrane does not bend if there is not a pressure difference between the two sides). On the other hand, a curved membrane exerts an overpressure on its convex side (conversely, a membrane bends if there is a pressure difference between the two sides). This effect can be observed for example when one takes a pot of yoghurt from sea level high into the mountains (the cover is bent outward because of the overpressure inside the pot). Figure 3 illustrates this effect. It is noted that the pressure exerted by a membrane is inversely proportional to its radius of curvature (Young-Laplace Equation [6]).

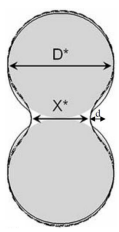


**Figure 3** A decrease in the radius of curvature of an elastic membrane requires an increased pressure. Source: MIT Non-Newtonian Fluid Dynamics Research Group.

If we consider two spherical grains that are in contact during a sintering process (Figure 4), we see that at the surface of the two grains the radius of curvature D/2 is positive while at the bridge connection between the two spheres the radius of curvature - d/2 is negative. The pressure difference **P** (normalized by the nominal external pressure **P0**) is proportional to the inverse of the radii of curvature of the surface of the grain and the connection area respectively (see equation (6)).

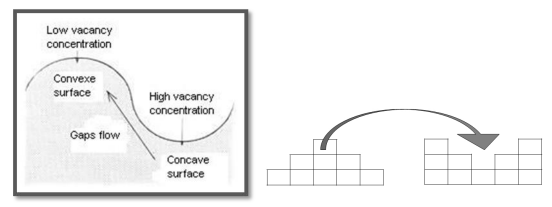
$$\frac{\Delta P}{P_o} \propto D^{*-1} + d^{-1} \tag{6}$$

This overpressure generates differences in the concentration of point defects and causes material transport to the bridge connection and thus the sintering of two spheres.



**Figure 4** Schematic representation of the shape evolution of two spheres in contact during a sintering process. D\* and d represent the diameters of curvature in two different local areas of the overall. Source: Yu U. Wang [11].

At the atomic scale, we can consider the number of neighboring atoms on a concave or convex surface. A concave surface has a higher average number of neighboring atoms than a convex surface; it follows that the atoms of a convex surface have a higher energy and greater mobility than those of a concave surface. Consequently grain surfaces act as sources and connecting bridges as sinks of material. Figure 5 shows a schematic view of the situation in two dimensions.



**Figure 5** Schematic of mass transport from a convex to a concave surface and an illustration of the number of neighboring atoms at a convex and concave surface respectively in two dimensions. Source: [6]

There are also other "sources" of matter within the green body during sintering, including atoms located at grain boundaries and defects within the material (e.g. dislocations). These sources become important as soon as there is a percolation of the green body (the time when one begins to have a continuous 3D solid structure), as from that moment, the migration of surface atoms will only change the shape of the pores (and not reduce their volume). Therefore, at this stage of sintering, the

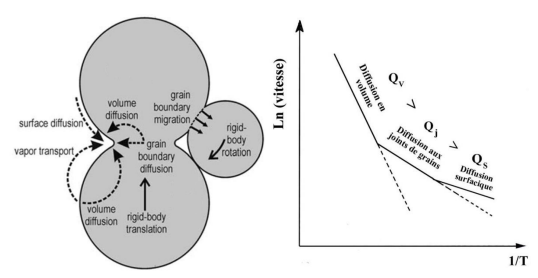
contribution of the surfaces as a source of material for densification is minimal, and it is the migration of atoms from grain boundaries and intra-granular defects that will be responsible for further densification. The driving forces causing matter transport from the grain boundaries and from the intra-granular defects are the same as for the surfaces: at the local level the negative radius of curvature at the connecting bridges, and globally the reduction of the interfacial energy will cause the matter transport.

For grain growth the major driving force is the reduction of the grain boundary energy. Indeed, the larger grains have a larger radius of curvature and a smaller interface to volume ratio and consequently a smaller energy, causing their growth at the expense of the smaller grains.

#### 3.2. Mechanisms of natural sintering

There are different sintering mechanisms (Figure 6), or in other words, different modes of matter transport from sources (surfaces, grain boundaries, defects) to the sinks (bridge connection). The different mechanisms are:

- Surface diffusion
- Volume diffusion
- Vapor transport
- Grain boundary diffusion (intergranular diffusion)



**Figure 6** Sintering mechanisms and dependence of the dominant mechanism on the sintering temperature.

Source: [6]

Not all the mechanisms are active for all sources of material. To transport material from intra-granular defects the only active mechanism is volume diffusion. For the transport of material from the interior of the grain boundaries both volume diffusion and grain boundary diffusion are active. To transport material from the surfaces, all mechanisms are active. For all the different modes of transportation we can define different diffusion coefficients  $\mathfrak D$  and their temperature dependence (Equation (7) [1]).

$$\mathcal{D} = \mathcal{D}_{o} e^{-\frac{Q}{kT}} \tag{7}$$

Where Q is the experimentally measured activation energy.

If we suppose for example that we have a vacancy diffusion mechanism (see Figure 7) we can define the diffusion coefficient depending on the vacancy concentration  $\mathbf{c}$ , the number of neighboring sites  $\gamma$ , the frequency of vibration of atoms  $\mathbf{v}$ , the distance between neighboring sites  $\lambda$  and the activation energy for vacancy jump  $\Delta \mathbf{G}$  (equation (8) [1]).

$$\mathcal{D} = c\gamma^{-1}v\lambda^{2}e^{-\frac{\Delta G}{RT}} = c\gamma^{-1}v\lambda^{2}e^{\frac{\Delta S}{R}}e^{-\frac{\Delta H}{RT}}$$
(8)

The only term of this equation that depends explicitly on the temperature is  $e^{\Delta H/RT}$  and thus at first approximation, the experimentally measured activation energy corresponds to the enthalpy of a vacancy jump  $\Delta H$ .

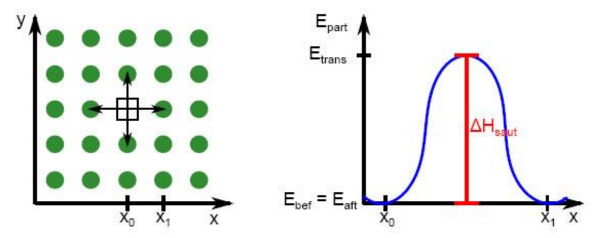


Figure 7 Schematic view of the vacancy diffusion and the activation energy for the vacancy diffusion.

In reality, there are other parameters, such as the concentration of vacancies c, which varies with the temperature and therefore influences the experimentally measured activation energy Q. This means that Q corresponds only approximately to  $\Delta H$ .

As the crystalline structure is disturbed at the interfaces of the grains (surfaces, grain boundaries), the activation energies for diffusion are most of the smaller solid interfaces than for scattering into volume ( $\Delta H_{surf} < \Delta H_{grain \, b.} < \Delta H_{volume}$  and therefore  $Q_{surf} < Q_{grain \, b.} < Q_{volume}$  see Table 1).

**Table 1** Diffusion coefficients for different diffusion mechanisms in  $\alpha$ -alumina and ZnO.

	$\alpha - Al_2O_3$ [2]	ZnO [3, 4]
Q <sub>surf</sub> [kJ/mol] Q <sub>grain b.</sub> [kJ/mol] Q <sub>volume</sub> [kJ/mol]	399 ± 122 464 ± 49 616 ± 102	$\sim 158$ $282 \pm 46$ $376 \pm 68$

There is often a diffusion mechanism that dominates the densification process. Thus the variation of the densification rate  $\rho^*$  with the temperature depends on the activation energy  $\mathbf{Q}$  of dominant diffusion mechanism. Because of the differences between the activation energies it is possible to determine which diffusion mechanism is dominant by measuring the density evolution under different conditions (e.g. different heating rates) using a dilatometer. A dilatometer measures the change of one of dimensions of the sample during sintering. Assuming that the densification is isotropic, we can then calculate the density  $\rho$  and the densification rate  $\rho^*$  from the change of dimension  $\mathbf{L}$  observed (equations (9) & (10)).

Practical classes

$$\rho = \frac{\rho_o}{\left(1 - \left|\frac{\Delta L}{L_o}\right|\right)^3} \tag{9}$$

$$\dot{\rho} = \frac{d\rho}{dt} = -\frac{3\rho_o}{\left(1 - \left|\frac{\Delta L}{L_o}\right|\right)^4} \frac{dL}{dt}$$
 (10)

It has been shown by several authors [7, 10] that one can separate the influences of temperature, density and grain size on the densification rate (equation (11)):

$$\rho = A \frac{e^{-\frac{Q}{RT}}}{T} \frac{f(\rho)}{D^n}$$
 (11)

Where **A** is a material parameter,  $\mathbf{f}(\rho)$  is a non specified function which depends only on  $\rho$  and  $\mathbf{n}$  depends on the dominant mechanism. In other words, if we measure the densification rate at different temperatures and roughly equal densities and grain sizes, we can estimate the activation energy by measuring the slope of  $\mathbf{ln}$  ( $\mathbf{T}$   $\rho$ ) versus  $\mathbf{T}^{-1}$  (equation (12)).

$$\ln(T\rho)\Big|_{\rho,D=cst} \propto -\frac{Q}{RT}$$
 (12)

Apart from the material transport by diffusion there are also stresses that develop during sintering and thus a movement of grains that helps densification.

The main mechanism responsible for grain growth is the movement of grain boundaries (in normal direction to the grain boundary) by local rearrangement of atoms. Indeed, at the grain boundaries there is the same phase at either side of the interface, no material transport is necessary for their movement. It is "sufficient" that the crystal structure near the grain boundary changes to be conform with the crystallographic orientation of the other grain and thus the grain boundary advances. However the situation changes if there are pores at the grain boundaries. As the energy of the pores is generally smaller at the grain boundaries than within the grains they often move with the grain boundaries. The movement of the pores is controlled by surface diffusion, a process much slower than the movement of grain boundaries by local rearrangement of atoms.

#### 3.3. Important parameters influencing the sintering

#### Grain size distribution and shape

The characteristics of the grains, that is to say the particle shape, size distribution of average grain size as well as the factor of agglomeration of the initial powder, strongly influences the sintering behaviour. The driving force and the kinetics are influenced by particle size. The total driving force for the sintering is proportional to the total surface energy per volume ( $\mathbf{F}_{sur} f^{N}$ ) incorporated in the

material.  $\mathbf{E}_{sur f}/V$  decreases with increasing grain size and when the shape of the grains approaches the equilibrium morphology (Wulff shape).

The densification kinetics is mainly influenced by variations in distances between sources and sinks of

material. For example, an irregular shaped grains or a large factor of agglomeration decreases the initial density of the green body, increasing the distance between the sources and the sinks. Consequently the rate of densification decreases. In addition the dominant diffusion mechanism can change with particle size and/or shape: the higher the specific interface area, the more the interfacial diffusion mechanisms (surface diffusion, grain boundary diffusion through the grain boundaries and vapor transport) are favored compared to volume diffusion.

The kinetics of grain growth is mainly influenced by the grain size distribution. If the distribution is large, the pressure difference between the smaller and larger grains is very high and consequently the growth of larger grains at the expense of the smaller ones is much faster than for a narrow size distribution.

#### **Temperature**

The temperature generally increases the rate of all sintering mechanisms. Also, as the activation energy (Q) is not the same for each diffusion mechanism, the rates of the different mechanisms might vary relatively to the rates of the other mechanisms. As in general  $Q_{\text{surf}} < Q_{\text{grain b.}} < Q_{\text{volume}}$  a higher temperature accelerates volume diffusion compared to interfacial diffusion. Thus the dominant densification mechanism may change with temperature. In addition, since grain growth is often controlled by surface diffusion while densification is controlled either by volume diffusion or by grain boundaries diffusion, a higher temperature often leads to higher densification compared to grain growth.

#### Dopants and impurities

There are often impurities and dopants present in the material. Dopants are added for various reasons such as changing the electronic properties (e.g.  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> doped with **Ti**), to facilitate sintering (e.g.  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> doped with **Mg**), or to improve the mechanical properties of the final product (e.g.  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> doped with **Y**). These dopants and impurities can affect the sintering in different ways:

- Formation of precipitates: If the concentration of impurities or dopants exceeds the solubility limit, a second phase will precipitate. As in the case of pores, the precipitates have an smaller energy at the grain boundaries than within the grains. Consequently second phase precipitates can decrease the mobility of grain boundaries and slow down grain growth.
- Increase of the concentration of vacancies or interstitial atoms: If the impurities / dopants have a different charge than the ions of the crystalline lattice there are often charged point defects such as vacancies or interstitial atoms that are created to compensate for the missing or excess charge. For example, in α- Al<sub>2</sub>O<sub>3</sub> doped with Mg, the magnesium ions have a charge of +2 while the replaced aluminum ion in the crystal have a charge of +3. The charges thus created are compensated by oxygen vacancies:

$$2MgO \stackrel{A12O3}{>} 2Mg'Al + V'O$$
.

Therefore dopants and impurities can influence the rate of diffusion by changing the concentrations of point defects.

- Immobilization of defects: the incorporation of a foreign ion in a crystal leads to the creation of a stress field around the defect. Bringing another defect with an inverse constraint field closer can therefore reduce their energy. For example the incorporation of Y³+ ions, which have a much larger ionic radius (0.9 Å) than Al³+ ions (0.54 Å), in α-Al₂O₃ creates a compression field around the Y. A vacancy on the other hand is surrounded by a tension field. Thus the total excess energy due to the presence of the two defects decreases when they approach each other. The resulting combination of an Y ion and a vacancy reduces the mobility of the two defects and thus dopant ions can decrease the diffusion coefficient.
- Segregation to interfaces: many dopants can be incorporated more easily at the interfaces of the material where the crystalline structure is already disturbed. Therefore dopants often segregate to interfaces. Similarly to pores and precipitates, the presence of segregated dopant ions at interfaces will decrease the mobility of grain boundaries as the dopants must migrate with the grain boundaries to maintain their low energy. In addition, the effects of eventual changes of the concentrations of point defects or of immobilizing the defects will be more pronounced at the interface due to the higher dopant concentration.

In summary, the exact effect of a dopant or impurity is complex, difficult to predict and depends on the ion type and concentration.

#### 3.4. Simulation of Sintering

Because of the complexity of sintering, it is often useful to use numerical simulations to estimate the influence of different factors on sintering. Therefore we will use a two-dimensional sintering model to simulate the influence of shape of the grains. The simulation is done on a cubic grid. Each grid cell can be occupied by a solid particle. Each particle has eight possible neighbors (see Figure 8). A number of neighboring solid particles can form a grain. The energy of a particle  $\mathbf{E}_{part}$  depends on the number of neighbors belonging to the same grain  $\mathbf{n}_{grain}$  and on the number of neighbors belonging to another grain  $\mathbf{n}_{boundary}$  (Equation 13).

$$E_{part=} n_{grain} \cdot \Delta E_{grain} + n_{boundary} \cdot \Delta E_{boundary}$$
 (13)

Where energies are adimensionalized:  $E^{adim} = \Delta H/RT$ . If at least one neighboring sites is not occupied, the particle can move. To move the particle must pass a transient state with a higher energy  $E_{trans}$  than the starting energy (see Figure 7). The average time the particle needs to move to a nearby site depends on the activation energy  $\Delta H_{saut} = (E_{trans} - E_{bef}) *RT$  of the jump.

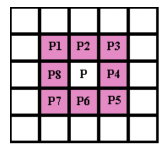


Figure 8 Neighborhood of a solid particle

Indeed time needed for a jump follows an exponential distribution that depends the activation energy (equation 14).

$$\rho(\Delta t) = (E_{trans} - E_{bef})e^{-(E_{trans} - E_{bef})\Delta t} = \frac{\Delta H_{saut}}{RT}e^{-\frac{\Delta H_{saut}}{RT}\Delta t}$$
(14)

Where  $\mathbf{E_{bef}}$  and  $\mathbf{E_{aft}}$  are the energies before and after the particle jump. In addition, if the particle has neighbors who do not belong to the same grain it can change the grain. The time distribution for a change from one grain to another is the same as to move from one site to another (equation 14). If the solid particle has no neighbors, and is thus in the vacuum, it behaves like a random walker with a constant speed  $\mathbf{V_{vaccume}}$ .

#### 3.5. Simulation of Y dopants in a-alumina

To estimate the influence of dopants on sintering we must use a model with atomic resolution. For example, to estimate the driving force for the segregation of dopants to interfaces, a model based on a pair potential can be employed, which enables the calculation of the energy contained in a specific structure. The analytical form of the pair potentials is composed of an electrostatic part (Coulomb) and an short-range interaction describing the interaction of electron clouds (Van der Waals) (see equation 15).

$$V_{ij}^{coulomb+van \text{ der Waals}} = \frac{q_i q_j}{4\pi\varepsilon_0 r_{ii}^2} + A_{ij} e^{-\frac{r_{ij}}{P_{ij}}} - C_{ij} r_{ij}^{-6}$$
(15)

Where A,  $\rho$  and C are empirical parameters determined by fitting the calculated properties of alumina and YAG (Y<sub>3</sub>Al<sub>5</sub>O<sub>12</sub>) (such as the Young's modulus or the crystalline unit cell dimensions) to the experimentally observed [13]. Summing over all pairs of ions, we can obtain the total energy contained in a specific structure. As the exact structure is often unknown, we try to minimize the energy by adjusting the position of the ions to obtain an equilibrated structure. Thus we can compare the energy of a dopant within the grains with the energy of a dopant at a surface or a grain boundary to calculate the driving force of segregation  $\Delta H_{seg}$  (equation 11).

$$\Delta H_{\text{seg}} = E_{\text{doped interface}} - E_{\text{pure interface}} - E_{\text{dopant}}$$
 (16)

Where  $\mathbf{E}_{doped\ interface}$  is the total energy of the interface containing the dopant,  $\mathbf{E}_{pure\ interface}$  is the total energy of the interface without dopants and  $\mathbf{E}_{dopant}$  is the energy of the dopant within the grains. There are several programs that were developed for this kind of calculations such as **METADISE** [9] and **GULP** [5].

### 4. Objectives

The objective of this lab is to understand the driving forces and mechanisms of natural, solid phase sintering and to develope an understanding of how they can be influenced. Sintering will be studied by different techniques. Firstly the Archimedes method will be used to study the porosity of samples sintered at different temperatures. Then experimental dilatometer results will be used to calculate activation energies. Thirdly numerical simulations of sintering with differently shaped grains will be

performed. In addition we will look at the behavior of Yttrium dopants in  $\alpha$ -Al<sub>2</sub>O<sub>3</sub>. The energy of the Y within the grains of  $\alpha$ -Al<sub>2</sub>O<sub>3</sub>, calculated with an atomistic model, will be compared with the energy of the Y at the  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> interfaces. The simulation results will also be compared to experimental observations. Finally some ideas how to incorporate the results of atomistic simulation into the sintering simulations will be developed.

#### 5. Work

- Measurement of the density of 3 sintered samples by the Archimedes method.
- Discussion of the densities (absolute and relative) as well as the open and closed porosity fractions of the three samples sintered at different temperatures.
- Launch of two simulations (with spherical and irregularly shaped grains). It is important to note
  - input parameters of the simulation and to periodically save an image of the current microstructure.
- Calculation of the activation energy using experimental dilatometer results for different heating rates. Comparison with the energies used for the sintering simulation.
- Calculation of the segregation energy of Y in Al<sub>2</sub>O<sub>3</sub> using atomistic simulations. Comparison with experimental observations (micrographs). How could the effect of dopants be included in the sintering model?

# 6. Requirements

### 6.1. Structure of report

The report must be structured as a scientific text, i.e. the content should be divided into several chapters that contain:

- A brief introduction that provides context and clarifies the objectives of the work done.
- The theoretical basis used for calculations and discussion.
- A description of the methods used.
- Results.
- A discussion of results.
- Conclusion.

Some of the points above can be combined into one chapter if preferred (e.g. Results and Discussion or Introduction and Theory ...).

#### 6.2. Content of report

The report should contain the following points:

- Effective density, percentage of open and closed porosity calculated for the three samples sintered at different temperatures.
- Description of the sintering model. For which such a model can be useful? What are the biggest limitations of the model?
- Calculation of the activation energy using experimental results for different heating rates. Comparison with the energies used for the sintering simulation.
- Results (microstructures, energy graphs, grain boundaries and surfaces based on the number of time steps) and discussion of the sintering simulation of varying particle size (spherical and irregular shaped grains). What are the phenomena observed? How the initial shape of the grains can influence the results? Comparison with experimental results for different particle sizes? How the porosity changes during sintering?
- Results of atomistic simulation ( $\Delta H_{\text{seg}}$  vs depth for the grain boundary and surface) and experimental observations (micrographs) for an  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> doped with Y. Discussion of results. How dopants could be included in the model of sintering?

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## Annex - Density of Water (g/mL) vs. Temperature (°C)

(from <u>Handbook of Chemistry and Physics</u>, 53rd Edition, p. F4) Whole **degrees** are listed down the left hand side of the table, while **tenths of a degree** are listed across the top. So to find the density of water at say **5.4** °C, you would first find the whole degree by searching down the left hand column until you reach '5'. Then you would slide across that row until you reach the column labeled '0.4'. The density of water at **5.4** °C is 0.999957 g/mL.

	0.0	0.1	0.2	0.3	0.4	0.5	0.6	0.7	0.8	0.9
0	0.999841	0.999847	0.999854	0.999860	0.999866	0.999872	0.999878	0.999884	0.999889	0.999895
1	0.999900	0.999905	0.999909			0.999923		0.999930	0.999934	0.999938
2	0.999941	0.999944	0.999947			0.999955		0.999960	0.999962	0.999964
3	0.999965	0.999967	0.999968			0.999971		0.999972	0.999973	0.999973
4	0.999973	0.999973	0.999973	0.999972	0.999972	0.999972	0.999970	0.999969	0.999968	0.999966
5	0.999965	0.999963	0.999961	0.999959	0.999957	0.999955	0.999952	0.999950	0.999947	0.999944
6	0.999941	0.999938	0.999935	0.999931	0.999927	0.999924	0.999920	0.999916	0.999911	0.999907
7	0.999902	0.999898	0.999893	0.999888	0.999883	0.999877	0.999872	0.999866	0.999861	0.999855
8	0.999849	0.999843	0.999837	0.999830	0.999824	0.999817	0.999810	0.999803	0.999796	0.999789
9	0.999781	0.999774	0.999766	0.999758	0.999751	0.999742	0.999734	0.999726	0.999717	0.999709
10	0.999700	0.999691	0.999682	0.999673	0.999664	0.999654	0.999645	0.999635	0.999625	0.999615
11	0.999605	0.999595	0.999585	0.999574	0.999564	0.999553	0.999542	0.999531	0.999520	0.999509
12	0.999498	0.999486	0.999475	0.999463	0.999451	0.999439	0.999427	0.999415	0.999402	0.999390
13	0.999377	0.999364	0.999352	0.999339	0.999326	0.999312	0.999299	0.999285	0.999272	0.999258
14	0.999244	0.999230	0.999216	0.999202	0.999188	0.999173	0.999159	0.999144	0.999129	0.999114
15	0.999099	0.999084	0.999069	0.999054	0.999038	0.999023	0.999007	0.998991	0.998975	0.998959
16	0.998943	0.998926	0.998910	0.998893	0.998877	0.998860	0.998843	0.998826	0.998809	0.998792
17	0.998774	0.998757	0.998739	0.998722	0.998704	0.998686	0.998668	0.998650	0.998632	0.998613
18	0.998595	0.998576	0.998558	0.998539	0.998520	0.998501	0.998482	0.998463	0.998444	0.998424
19	0.998405	0.998385	0.998365	0.998345	0.998325	0.998305	0.998285	0.998265	0.998244	0.998224
20	0.998203	0.998183	0.998162	0.998141	0.998120	0.998099	0.998078	0.998056	0.998035	0.998013
21	0.997992	0.997970	0.997948	0.997926	0.997904	0.997882	0.997860	0.997837	0.997815	0.997792
22	0.997770	0.997747	0.997724	0.997701	0.997678	0.997655	0.997632	0.997608	0.997585	0.997561
23	0.997538	0.997514	0.997490	0.997466	0.997442	0.997418	0.997394	0.997369	0.997345	0.997320
24	0.997296	0.997271	0.997246	0.997221	0.997196	0.997171	0.997146	0.997120	0.997095	0.997069
25	0.997044	0.997018	0.996992	0.996967	0.996941	0.996914	0.996888	0.996862	0.996836	0.996809
26	0.996783	0.996756	0.996729	0.996703	0.996676	0.996649	0.996621	0.996594	0.996567	0.996540
27	0.996512	0.996485	0.996457	0.996429	0.996401	0.996373	0.996345	0.996317	0.996289	0.996261
28	0.996232	0.996204	0.996175	0.996147	0.996118	0.996089	0.996060	0.996031	0.996002	0.995973
29	0.995944	0.995914	0.995885	0.995855	0.995826	0.995796	0.995766	0.995736	0.995706	0.995676
30	0.995646	0.995616	0.995586	0.995555	0.995525	0.995494	0.995464	0.995433	0.995402	0.995371

(Updated by C.R. Snelling, 6/14/08)