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2015 IOP Conf. Ser.: Mater. Sci. Eng. 84 012102
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Two-phase modelling of equiaxed crystal sedimentation and thermomechanic stress development in the sedimented packed bed

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Abstract. During many industrial solidification processes equiaxed crystals form, grow and move. When those crystals are small they are carried by the melt, whereas when getting larger they sediment. As long as the volume fraction of crystals is below the packing limit, they are able to move relatively free. Crystals being backed in a so called packed bed form a semi-solid slurry, which may behave like a visco-plastic material. In addition, cooling-induced density increase of both, liquid and solid phases might lead to shrinkage of the whole casting domain. So deformation happens and gaps between casting and mold occur. In the present work, a two-phase Eulerian-Eulerian volume-averaging model for describing the motion of equiaxed crystals in the melt is combined with a similar two-phase model for describing the dynamic of the packed bed. As constitutive equation for the solid skeleton in the packed bed Norton-Hoff law is applied. Shrinkage induced by density changes in the liquid or the solid phase is explicitly taken into account and handled by remeshing the calculation domain accordingly.

1. Introduction
In many practical solidification processes the equiaxed crystals that form first move with the melt when they are small and then sink down and sediment when they are getting larger. Specially, in big steel ingots or in grain refined Al-alloy slabs, large regions usually form by growth and sedimentation of equiaxed crystals. While the melt cools and crystals occur, liquid and solid densities changes with temperature. In addition, often there exist a density change between solid and liquid. For shape and ingot casting or even continuous casting, temperature and phase changed-induced density changes leads to massive shape deformations including the formation of shrinkage cavities and/or microporosity. In last decades, numerical simulation of nucleation, growth, motion and sedimentation of equiaxed crystals have made huge progress especially by means of multi-phase volume averaging methods [1-3]. Also, the simulation of deformation and stresses caused by uneven cooling from solidus to room temperature is a standard procedure today. However, the simulation of flow related phenomena on one side combined with stress and deformation which evolve already during solidification deep inside the mushy zone on the other side is still missing. This paper is supposed to do first steps to close this gap.
In the chemical industry a stack of small objects in a vessel is called packed bed. Much knowledge about permeability of dendritic mushy regions are gained from flow experiments with packed beds. Also the rheological behavior of packed beds made of porous metallic materials saturated with liquid at high temperatures were subject of investigations. Nguyen et al. found for Sn-Pb alloys in the semi-solid state that two functions are needed to describe the corresponding viscoplastic potential [4]. Bellet et al. [5,6] used this viscoplastic potential for the volume-averaged version of the effective macroscopic stress tensor of the solid phase in the mushy zone of a continuous casting of steel. In the present paper, we suggest to combine this approach with a two-phase Eulerian-Eulerian model for motion and sedimentation of equiaxed crystals and a method to deform the mesh in order to account for temperature-induced density changes.

2. Model description
2.1 Basic considerations
Based on former work of the authors [1-3], a two-phase Eulerian-Eulerian model for simulating formation and sedimentation of equiaxed crystals has been adopted and combined with the two-phase thermomechanic model suggested by Bellet et al [5-6]. The following main considerations were made:

Two phases are defined: the liquid phase (ℓ) and the solid (s). The corresponding phase fractions, g_ℓ and g_s, add up to one. A predefined number density of already existing grains was considered. The implementation of nucleation and growth of equiaxed crystals is under construction and will be topic of a subsequent paper. Mass-, momentum and enthalpy conservation is solved for both phases by considering (i) above the packing limit a submerged object drag law [6] and an effective solid viscosity which ensures the proper mixture behavior (Eq. (6),(8),(9)), and (ii) below the packing limit a Darcy term drag law with a Kozeny-Carman type permeability and a viscoplastic materials law for the packed bed (Eq. (6),(10)). Note that following the arguments in [6-7], the pressure equilibrium in the liquid phase is almost instantaneous. Subsequently, the interfacial pressures [8] in both phases equal the intrinsic average value of liquid pressure, i.e. its microscopic value.

We also allow that the density of the liquid and solid may vary with temperature. Thus, we have considered corresponding thermal strain rates according to [9]. For the packed bed regime, we use a volume-averaged Norton-Hoff law. As stated in [10], for metallic alloys at elevated temperature the constitution of the solid phase can be characterized by the Norton-Hoff law. According to [5-6] volume-averaging of this law gives a compressible viscoplastic constitutive model which tends to the incompressible viscoplastic Norton–Hoff model as solidification completes (Eq. (10)). As viscoplastic consistency, K_v, and strain rate sensitivity coefficient, m, we temperature-averaged the information given in [11] for the low carbon steel (with a composition in mass pct. of 0.06C, 0.1Ni, 0.13Mn, 0.1Si, 0.08Cu, 0.035Al, 0.015P, and 0.012S) and got m = 0.139. In the current studies we were not satisfied with the K_v value based on [11], predicting rather low effective viscosity at the viscoplastic regime. To derive the value of K_v, the assumption of the equality of viscous stress prefactors in Newtonian regime (Eq. (9)) and in viscoplastic one (Eq. (10)) at the coherence limit was made. By ignoring the strain rate influence (assuming m = 1) at the transition point, one can derive 2μ = 3K_v/μ_s A. From this algebraic calculation, we got K_v = 7668 at g_s = g_s,pack ≈ 0.637. Please note that the transition point is somehow arbitrary. With a transition at g_s = 0.55 K_v would have been much smaller and so the apparent viscosity of the viscoplastic semi-solid mush. It will be subject of future work to adapt the transition point to a suitable experimental measured K_v.
Table 1. List of equations solved on a grid with moving boundaries. Symbols are defined in the Nomenclature at the end of this paper.

mass conservation

\[
\frac{\partial}{\partial t} (g_s \rho_s) + \nabla \cdot (g_s \rho_s \mathbf{v}_s) = 0
\]

(1)

\[
\frac{\partial}{\partial t} (g_s \rho_s \mathbf{v}_s) + \nabla \cdot (g_s \rho_s \mathbf{v}_s \mathbf{v}_s) = \nabla \cdot (g_s \mathbf{\sigma}_s) + g_s \rho_s \mathbf{g} - K_s (\mathbf{v}_s - \mathbf{v}_s^{eq})
\]

(2)

momentum conservation

\[
\frac{\partial}{\partial t} (g_s \rho_s \mathbf{v}_s) + \nabla \cdot (g_s \rho_s \mathbf{v}_s \mathbf{v}_s) = \nabla \cdot (g_s \mathbf{\sigma}_s) + g_s \rho_s \mathbf{g} + K_s (\mathbf{v}_s - \mathbf{v}_s^{eq})
\]

(3)

\[
K_s = 18g_s \mu_s \frac{g_s C_v}{d_s^2} \quad \text{for} \quad g_s < g_{s, pack}
\]

(4)

\[
= g_s \frac{\mu_s}{K} \quad \text{for} \quad g_s \geq g_{s, pack}
\]

with \( K = d_s^2 \frac{1}{180 g_s^3} \).

\[
\mathbf{\sigma}_s = -p_s \mathbf{I} + \mathbf{\tau}_s, \quad \mathbf{\tau}_s = 2\mu_s \left( \dot{\mathbf{\epsilon}}_s - \frac{1}{3} \text{tr}(\dot{\mathbf{\epsilon}}_s) \mathbf{I} \right), \quad \dot{\mathbf{\epsilon}}_s = \frac{1}{2} \left( \nabla \mathbf{v}_s + \left( \nabla \mathbf{v}_s \right)^T \right), \quad \text{tr}(\dot{\mathbf{\epsilon}}_s) = \nabla \cdot \mathbf{v}_s
\]

(5)

\[
\text{for} \quad g_s < g_{s, pack}:
\]

\[
g_s \mathbf{\sigma}_s = 3K_s \left( \sqrt{3} \dot{\mathbf{\epsilon}}_{eq}^* \right)^{m-1} \left( \frac{1}{A} \dot{\mathbf{\epsilon}}_s - \frac{1}{A} \left( \frac{1}{3} \text{tr}(\dot{\mathbf{\epsilon}}_s) \mathbf{I} \right) \right)^{m-1}
\]

(6)

with \( \dot{\mathbf{\epsilon}}_{eq}^* = \frac{1}{A} \sum_{i,j} \dot{\mathbf{\epsilon}}_{s,ij} \dot{\mathbf{\epsilon}}_{s,ij} - \frac{3}{A} \left( \frac{1}{B} \text{tr}(\dot{\mathbf{\epsilon}}_s) \right)^2 \) and defining \( \eta_s^{opp} = \frac{3K_s}{g_s \rho_s A} \left( \sqrt{3} \dot{\mathbf{\epsilon}}_{eq}^* \right)^{m-1} \)

enthalpy conservation

\[
\frac{\partial}{\partial t} (g_s \rho_s h_s) + \nabla \cdot (g_s \rho_s \mathbf{v}_s h_s) = \nabla \cdot (g_s \mathbf{k}_s \nabla T_s) + H^* (T_s - T_s)
\]

(7)

\[
\frac{\partial}{\partial t} (g_s \rho_s h_s) + \nabla \cdot (g_s \rho_s \mathbf{v}_s h_s) = \nabla \cdot (g_s \mathbf{k}_s \nabla T_s) - H^* (T_s - T_s)
\]

(8)

\[
h_s = \int_{T_{ref}}^{T_s} c_{pl(s)}dT + h_{s,ref}^T \quad \text{and} \quad h_s = \int_{T_{ref}}^{T_s} c_{pl(s)}dT + h_{s,ref}^T \quad \text{and} \quad H^* = 10^8 \text{ W/m}^3\text{K}
\]

(9)

\[
\text{grain number density conservation and average grain diameter}
\]

\[
\frac{\partial}{\partial t} (n_s) + \nabla \cdot (n_s \mathbf{v}_s) = 0
\]

(10)

\[
d = 2 \sqrt{\frac{3g_s}{4\pi n_s}}
\]

(11)
The two rheological functions $A(g_s)$ and $B(g_s)$ used in Eq. (10) are valid for an equiaxed dendritic network (backed bed) and were taken from [4] and reprinted in Table 2. They satisfy $A(1) = 3/2$ and $B(1) = 0$, imposing in that way the incompressibility constraint by means of a penalty factor, the multiplier of $\dot{\varepsilon}$ in Eq. (10), which becomes infinite. Thus, the second term of Eq. (10) vanishes and the equations turns into the regular Norton-Hoff law. Numerically, the penalty factor is made arbitrarily high but finite by replacing $B(1)$ by $B(0.999)$. As coherency limit we have chosen $g_{s,cohe} = g_{s,pack}$.

All equations solved are gathered in Table 1.

2.2. Numerical implementation

The numerical results are obtained by performing the two phase simulation using the in-house multiphase solver developed in the open-source CFD package OpenFOAM®. The heat transfer is included to track the domain shrinkage due to the temperature dependent density change applying the numerical grid motion / smoothing technique.

Table 2. Rheological function $A(g_s)$ and $B(g_s)$ according to Table 1 in [4]

<table>
<thead>
<tr>
<th>$A(g_s)$</th>
<th>$B(g_s)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$3/2 \cdot g_s^{0.47}$ for $g_s &gt; g_{s,cohe}$</td>
<td>$0.009 \cdot \left[\frac{1}{g_s^{0.94}} - 1\right]$ for $g_s &gt; g_{s,cohe}$</td>
</tr>
</tbody>
</table>

The mesh motion algorithm is based on the information of the mass fluxes at the domain boundaries required to compensate the volume loss. Therefor a compressible mixture sub-model is applied with the opened domain boundaries as opposed to the closed one for the multiphase model. The boundary displacements are calculated by interpolating the velocity at the face centers to the boundary points. After the boundary points are moved, the internal mesh is smoothed by solving a Laplace equation for the cell centers to maintain the mesh quality [12-14]. This procedure is done for each time step and the coupling between mesh motion and flow calculation is repeated until the zero boundary fluxes are reached.

The presented shrinkage model implementation consists of the following steps:
1. The solution of the multiphase model is obtained for the current iteration.
2. Updated density distribution is calculated.
3. Boundary mass fluxes are calculated from the additional compressible mixture model to compensate the density change, provided from the settling and the heat transfer simulation. Gravity is excluded from the calculations to prevent the liquid / solid mixture run-out.
4. The mesh is moved along the boundaries and sequentially smoothed using a Laplace equation with the provided mesh diffusion coefficient: the bigger coefficient is the more the deformation of the boundaries is smoothed and redistributed through the all mesh elements. Mesh motion fluxes are then included into the convective terms of the multiphase model.
5. The complete multiphase model is recalculated for the next iteration on the modified mesh.
6. Mesh motion / solution updates are continuously repeated until the shrinkage is compensated.
7. The solution is advanced for the next time step.
The simulation domain is shown in Fig. 1 and represents a square cavity (each side is 10 mm), which is isolated from all sides except for the bottom one. The bottom wall is cooled down to the reference cooling temperature 1173.15 K and the heat transfer coefficient of 1000 W/m²K.

The simulation starts with a uniform distribution of 70% liquid and 30% solid in the whole domain and the initial temperature being uniform for both phases. All walls of the cavity are nonslip and initially we start with a no-flow situation.

![Fig. 1 Domain with the boundary and initial conditions](image)

### Table 3. Material properties used in the test case

<table>
<thead>
<tr>
<th>Material</th>
<th>$T_{ref}$ (K)</th>
<th>$c_p$ (J/kg/K)</th>
<th>$k$ (W/m K)</th>
<th>$\rho$ (kg/m³)</th>
<th>$\alpha$ (1/K)</th>
<th>$d$ (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>liquid</td>
<td>1803.44</td>
<td>820.6</td>
<td>35.846</td>
<td>7021.95</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>solid</td>
<td>1803.44</td>
<td>735.9</td>
<td>35.771</td>
<td>7291</td>
<td>$63 \cdot 10^6$</td>
<td>$10^{-4}$</td>
</tr>
</tbody>
</table>

### 3. Results and discussion

Results of the simulation are shown in Fig. 2, where the evolution of the solid fraction distribution is presented. At the very beginning, both phases obey the Newtonian liquid behavior, and the phase separation interface is quite uniform (see Fig. 2a). Later on, the high concentration of the solid above the packing limit (marked with a black line at Fig. 2b) is reached at the bottom of the domain; the apparent (effective) viscosity of the solid phase rapidly increases following the visco-plastic rheology described above. The formation of the coherent solid structures with freckles in between (Fig. 2c) providing the passages for the lighter liquid can be seen. During their development the high solid content zones tend to collapse and to merge together, entrapping some liquid pockets.

The density change for the solid phase is based on the temperature distribution and the compressibility factor $\alpha$ (see Table 3). One can follow the deformation history of the cavity with time in Fig. 2a-f. It can be noted that (i) the total domain volume is reducing; and (ii) the bottom of the cavity has shrunk more due to the higher concentration of the solid phase, which is the only phase compressing in the presented studies. Please note that all deformations are multiplied by a factor of 2 to make them noticeable on the figures.
Fig. 2 Evolution of the solid fraction field. The dotted rectangle in (c) is analyzed further in Fig. 3. The arrow in (d) indicated the inner granular melt flow.

Fig. 3. Results at time $t = 50$ sec: solid fraction with (a) liquid velocity and (b) solid velocity vectors; (c) apparent kinematic viscosity of the solid $\mu_{\text{app}}$ distribution; (d) shrinkage rate: ratio of the calculated mixture density $\rho(T)$ related to the initial isothermal density $\rho_0$; isotherms show the mixture temperature distribution.
Fig. 4. Collapse of a single solid phase structure: (a) initial moment; solid velocities are shown as vectors; (b) collapsed structure is merged with the more stable one near the cavity wall. The velocity vectors in the large picture show the liquid velocity escaping from the mush.

With time the initially more or less uniform phase interface becomes quite distorted due to the formation of stable solid (sub-)structures. More other, some zones with higher liquid content (here called liquid pockets) are entrapped between the solid structures.

After the solid phase finally settles at the bottom of the domain, the rest of the liquid floats up through the channels (similar to one marked with a red arrow in Fig. 2d) before they are closed with the coherent solid zones (see Fig. 2c). The stresses occurring in the coherent solid structures are so strong, that they resist any pressure from the uprising liquid, and the rest of it remains entrapped and the final shape of the solid fraction field remains almost unchanged until the system cools down, as shown in Fig. 2e-f.

As it was already mentioned, the heat transfer is simulated together with the solid phase sedimentation causing local density change in the solid phase and the shrinkage of the domain as a consequence of that. Fig. 3 shows the details of the solid / liquid phases interaction at $t = 50\text{sec}$ at the bottom part of the simulation domain, marked with the red dashed rectangle in Fig. 2c. One can see that liquid tends to pass through the freckles / channels in the coherent solid structures, as shown with the vector field in Fig. 3a. Correspondingly, the solid phase sinks down and its motion is dominantly downwards (as one can see from Fig. 3b). However, due to the high viscoplastic stresses which appear above the packing limit, the solid velocities are much smaller if compared to the liquid ones (vector scaling is the same in Fig. 3a-b). The distribution of the apparent (effective) viscosity $\mu_{s}^{app}$ of the solid is plotted in Fig. 3d: its maximum value is 6÷9 orders higher than in the liquid, causing a strong resistance to the strain in the dense solid regions. Additionally the shrinkage ratio is presented in Fig. 3e corresponding to the increased density (due to the thermal compression in the domain) to the initial density relation.

Finally, an example of the coherent behavior in the highly packed solid is illustrated as an avalanche / collapsing of the single bound structure (see Fig. 4). The results are taken from an additional simulation not included in the current study. So, if we follow the motion of the marked object, its starts to fall down in the direction of a white arrow in Fig. 4 and to rotate simultaneously, after the small bridge (marked with the red dotted line) collapses under its weight and the liquid escapes from a freckle formed in between coherent solids. In Fig. 4a solid velocities are shown as a vector field indicating that the
bound solid structure moves as a single rotating object, until it impacts by its tip and merges with another more stable one close to the cavity wall.

4. Summary
In order to model both, motion and sedimentation of equiaxed crystals together with stress and deformation in the backed bed, a two-phase Eulerian-Eulerian volume-averaging model is suggested which uses different constitutive models for the stress-strain rate relation of the solid phase below and above the packing limit. In addition, shrinkage induced deformation of the whole domain is accounted for by an adequate remeshing model. As result, the formation of a backed bed of sedimenting equiaxed crystals is models together with the melt flow induced by the sedimentation process. As the solid skeleton is modelled by a visco-plastic flow model, also deformation of and stress evolution in the backed bed is predicted. The results show that the actual dynamic of sedimentation and visco-plastic and density-related deformation might be quite complex as semi-solid channels where liquid can flow more easily might exists in between coherent solid area. The actual dynamic of this process was found to depend highly on the assumed material parameter. As an example, the formation of a continuous crystal pill up and a corresponding initiation of a crystal avalanche / collapsing was shown. The model is the first step towards describing the formation of the solidifying shell together with the occurring gap between mold and metal during thin slap casting of steel.

Acknowledgement
This work was financially supported by the FWF Austrian Science Fund (P22614-N22), FFG Bridge Early Stage (No. 3893791), and the Austrian Federal Ministry of Economy, Family and Youth and the National Foundation for Research, Technology and Development within the framework of the Christian Doppler Laboratory for Advanced Process Simulation of Solidification and Melting.

Nomenclature

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>A, B</td>
<td>rheological function taken from [4]</td>
</tr>
<tr>
<td>( c_{p(s)} )</td>
<td>( \ell/s ) specific heat [J/kg/K]</td>
</tr>
<tr>
<td>( C_s )</td>
<td>settling ratio [-]</td>
</tr>
<tr>
<td>( d )</td>
<td>vol. averaged grain diameter [m]</td>
</tr>
<tr>
<td>( g )</td>
<td>gravity acceleration [m/s²]</td>
</tr>
<tr>
<td>( g_{s,pack} )</td>
<td>packing limit [-]</td>
</tr>
<tr>
<td>( g_{s,cohe} )</td>
<td>coherency limit [-]</td>
</tr>
<tr>
<td>( h_s, h_s' )</td>
<td>( \ell/s ) volume averaged enthalpy [J]</td>
</tr>
<tr>
<td>( h_s'^{ref}, h_s'^{ref} )</td>
<td>( \ell/s ) enthalpy at ( T_{ref} ) [J]</td>
</tr>
<tr>
<td>( I )</td>
<td>identity tensor [-]</td>
</tr>
<tr>
<td>( K )</td>
<td>much permeability [m²]</td>
</tr>
<tr>
<td>( K_v )</td>
<td>viscoplastic consistency [kg s⁻²/m]</td>
</tr>
<tr>
<td>( k_s )</td>
<td>( \ell/s ) eff. heat conductivity [J/m²K]</td>
</tr>
<tr>
<td>( m )</td>
<td>strain rate sensitivity coefficient [-]</td>
</tr>
<tr>
<td>( n_s )</td>
<td>grain number density [-]</td>
</tr>
<tr>
<td>( p_r )</td>
<td>intrinsic pressure in the liquid [kg/s²/m]</td>
</tr>
<tr>
<td>( \rho_s )</td>
<td>( \ell/s ) densities [kg/m³]</td>
</tr>
<tr>
<td>( \tau_s )</td>
<td>( \ell/s ) vol. av. dev. stress tensor [kg/s²/m]</td>
</tr>
<tr>
<td>( \mu_s )</td>
<td>( \ell/s ) viscosity [kg/m/s]</td>
</tr>
<tr>
<td>( \mu_s^{app} )</td>
<td>apparent kinematic viscosity of the solid</td>
</tr>
<tr>
<td>( \alpha )</td>
<td>compressibility [1/K]</td>
</tr>
</tbody>
</table>

References