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ON A MUSHY CELL TRACKING METHOD FOR SIMULATING GALLIUM MELTING

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agree well with the results from other numerical solutions. the boundary conditions at the interface of the mushy, solid, and liquid cells becomes feas-ible. The predicted moving interface and thermofluid field of gallium melting are shown to dritic solidification. mushy cells without consideration of the curvature and normal velocity effects in the denlies in the specification of constant melting or freezing temperature at the center of the located in the cell centers, the distinguishing characteristic of the present tracking algorithm under the mass and energy balance laws to capture the mushy cell front. With the variables tured grid using the finite-volume formulation. The mushy cell tracking equation is derived expressed in terms of the primitive variables and are discretized in the stationary unstrucmushy cell placed in between. The governing equations for the thermofluid transport are computational domain is separated into solid-phase and liquid-phase regions, with the cell tracking method to clarify whether the solution is physically correct. In the study, the converged solution remained, we reexamine this problem using the thermally driven mushy ied. Since the previously simulated results were not consistent and the myth of a grid-Gallium melting in a rectangular cavity heated from the side wall has been extensively stud-Thanks to this feature, a straightforward and accurate evaluation of

INTRODUCTION

can change sharply. Nonlinear physical complexities emerge from a wide range of some internal boundaries or interfaces, across which the physicchemical properties tational fluid dynamics scientists [7, 9, 14]. This class of problems is characterized by tered in natural environments and processing industries have been made by compu-In the past two decades, numerous studies of phase-change problems encoun-

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	NOMENCLATURE	TATU	J R E
A_f	outward area vector for the face _f	δt	time step
P	pressure	ν	kinetic viscosity
St	Stefan number, $C_s\Delta T/\Delta H$	ρ	density
t	time		gradient operator
u_i	velocity component along the i-direction	\mathfrak{Q}_0	domain of <i>cell</i> ₀
\overline{U}	velocity vector	$\ \Omega_0\ $	volume of Ω_0
\underline{U}_f	face velocity vector		

enthalpy-based [10, 11] and moving = particle [12] methods. Shyy [7] has given a employed for solving free/moving-boundary problems can be classified into the for this class of problems are only available for very few cases with simple geometry moving interface itself is considered as a part of the solution process for solving urations and bodies with complex shapes, can further complicate the analysis. detailed review of these methods. their geometric and physical complexities, among other factors. Analytic solutions Free-and moving-boundary problems are normally difficult to analyze because of time and space scales. In addition, geometric complexities, such as multibody configvarying interfaces has gradually become feasible. In the literature, the methods 14]. With the advent of high-speed computers, numerical capturing of the timesystem of equations governing the behavior of the thermofluid flow field.

considering curvature and normal velocity effects in the algorithm is the specification of constant melting or freezing temperature without various fields can be found in [5, 6]. The distinguished characteristic of the present energy balance principles. A detailed description of the method and its application to tational mushy cell, for the current two-phase problem is governed by the mass and region is governed by the stable diffusion process, while the transport phenomenon the mushy cell placed in between. The transport phenomenon in the solid-phase computational domain is separated into solid-phase and liquid-phase regions, with ithm [5, 6] is applied to simulate 1 D solidification and 2 D tin melting. After invesrequired. In this study, the cell-by-cell thermally driven mushy cell tracking algorof mass, momentum, and energy along with the tracking of the moving interface are of liquid and melting of solid, the working equations for describing the conservations distinguishing features [5]. comparison with other methods, the mushy cell tracking algorithm has the following diffusion process. The movement of the interface, which is referred to as the compuof the liquid-phase region is governed by the conditionally stable tigating these two benchmark tests, the gallium melting problem is investigated. The boundary conditions at the interface between the mushy, solid, and liquid cells. [22]. Thanks to this embedded feature, one can easily and accurately evaluate the When a numerical method is applied to solve problems involving solidification dendritic solidification convection-

- The volume-averaged property $\langle F \rangle_{m}$ is introduced.
- Solidification or melting time is defined as a function of $\langle F \rangle_m$
- Evaluation of $\partial T/\partial t$ in the mushy cells is circumvented.

cing any source term involving the volume-averaged $\langle F \rangle_m$ explicitly. The field variables for both liquid and solid phases are solved without introdu-

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4.

- S to a local point of the moving front. The size of the region of $\langle F \rangle_m$ is controlled by the mushy cell, and can be reduced
- 6 solid or liquid cells. temperature condition is specified at the interfaces between the mushy and either The algorithm is mainly controlled by a conduction-type equation, and there is no need to solve for the temperature in the mushy cells since a constant melting
- .7 the field variables are solved in a phase-by-phase and cell-by-cell manner. The mushy cells separate the domain into liquid and solid zones. Consequently,

with an aim of clarifying the afore-mentioned controversial problem. problem will be reexamined in this study using the mushy cell tracking algorithm melting problem and tried to resolve some previously known controversies. and numerically [2-4]. Hannoun et al. [4] gave an excellent review of the gallium numerical model. The gallium melting problem has been studied experimentally [1] known to be amenable to analytical solution, will be used to access the proposed is the so-called Stefan problem. The predicted interface location, which has been ation of convection effect [1–4, 11, 13, 15, 16] will be investigated. The first problem ation of convection [14] and the melting of tin in a rectangular cavity with consider-To validate the present algorithm, both the 1-D solidification without consider-

2. WORKING EQUATIONS

are expressed as dividing it into three distinct domains, Ω_s (solid zone), Ω_m (mushy zone), and Ω_L (liquid zone), where $\Omega_s \cup \Omega_m \cup \Omega_L = \Omega$. The flow and energy conservation equations this problem was first published by Stefan [14] to study the thickness of the polar ice This classical problem considers the energy conservation in the domain Ω by The solidification problem is sometimes referred to as the Stefan problem, since

$$\underline{\nabla} \bullet \underline{U} = 0 \quad \text{in } \Omega_L \tag{1a}$$

$$\rho_L \left[\frac{\partial \underline{U}}{\partial t} + \underline{\nabla} \bullet (\underline{U}\underline{U}) \right] = -\underline{\nabla}P + \underline{\nabla} \bullet \underline{\underline{\tau}} - \rho_L \underline{g} \beta_T (T - T_{\text{ref}}) \quad \text{in } \Omega_L$$
 (1b)

$$\rho_L C_P \left[\frac{\partial T}{\partial t} + \underline{\nabla} \bullet (\underline{U}T) \right] = \underline{\nabla} \bullet k_L \underline{\nabla}T \quad \text{in } \Omega_L$$
 (1c)

$$\rho_S C_P \frac{\partial T}{\partial t} = \underline{\nabla} \bullet k_S \underline{\nabla} T \qquad \text{in } \Omega_S$$
 (1*d*)

where

$$\equiv \mu \left[\left(\underline{\nabla U} \right) + \left(\underline{\nabla U} \right)^t \right] - \frac{2\mu}{3} \underline{\nabla} \bullet \underline{U} \quad \text{in } \Omega_L$$
 (2)

lla

moving-interface restriction. The internal boundary condition is specified as It should be noted that the current study is investigated under the isothermal

$$T = T_m \quad \text{in } \Omega_m(t) \cup \partial \Omega_m(t)$$
 (3)

mal expansion coefficient β_T . In Eq. (1), \underline{U} is the fluid velocity and P is the pressure eration \underline{g} , temperature T, dynamic viscosity μ , thermal conductivity k, and the ther-Other physical properties involved are the density ρ , specific heat C_P , gravity accel-The subscripts s, L, and m denote the solid, liquid, and mushy states, respectively

the Stefan condition [14] should be valid at the solid/liquid interface for truly repremushy cell. When the moving-particle method is used to trace the front propagation, equation is needed in order to account for the energy balance at the interface of the solid heat conduction. To describe the movement of the mushy cell, one additional tion, the equation for the thermal energy can be reduced to one similar to that for senting the solidification process, When considering phase-change problems without taking into account convec-

$$\rho_s(\Delta H)\underline{W} \bullet \underline{n} = k_s \underline{\nabla} T_s \bullet \underline{n} - k_L \underline{\nabla} T_L \bullet \underline{n}$$
(4)

tracking approach presented later on. the computational time and memory can be significantly reduced in the mushy cell the computational time and memory requirements can be excessive. However, both melting or frozen temperature criterion. When using this front-capturing method, of the moving front. It should be noted that Eq. (4) corresponds to the constant normal derivatives of T in the liquid and solid regions, respectively. \underline{W} is the velocity the outward normal vector of the moving front, and $\nabla T_L \bullet \underline{n}$ and $\nabla T_s \bullet \underline{n}$ denote the In the above equation, ΔH denotes the latent heat of solidification. The vector \underline{n} is

the conservation of mass and energy can be written in integral forms as follows: the nonmoving framework using the control-volume method. The equations for During solidification, the mushy cell tracking equation will be derived within

$$\frac{\partial}{\partial t} \int_{\Omega_m} \rho \ d\Lambda + \oint_{\partial \Omega_m} \rho(\underline{U} - \underline{W}_m) \bullet d\underline{A} = 0 \tag{5}$$

$$\frac{\partial}{\partial t} \int_{\Omega_m} \rho H \, d\Lambda + \oint_{\partial\Omega_m} \rho H (U - \underline{W}_m) \bullet d\underline{A} = \oint_{\partial\Omega_m} \underline{q} \bullet d\underline{A} \tag{6}$$

In the above equations, \underline{W}_m is the control-surface velocity, which is zero for a fixed control volume, Ω_m is the fixed control volume, and \underline{q} is the conduction heat flux. Here, the following two important assumptions will be made:

- treated as a porous medium with a fairly high resistance to the flow. No flow is allowed to proceed between the mushy cells. Actually, the mushy cell is
- Across the interface, no temperature gradient is allowed to take place because the temperature of each mushy cell is specified as T_m . Taking the mushy cell shown in

Figure 1 as an example, the mass and energy equations can be expressed as

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$$\|\Omega_m\|_{\widehat{\partial t}}^{\widehat{\partial}}\langle\rho\rangle_{\Omega_m} + \sum (\rho_L(\underline{U}_{mL} - \underline{0}) \bullet \underline{A}_{mL}) = 0$$
 (7)

$$\|\Omega_m\| \frac{\partial}{\partial t} \langle \rho H \rangle_{\Omega_m} + \sum_s (\rho_L H_{mL}(\underline{U}_{mL} - \underline{0}) \bullet \underline{A}_{mL})$$

$$= \sum_s (k_s \underline{\nabla} T_s \bullet \underline{A}_{ms}) + \sum_s (k_L \underline{\nabla} T_L \bullet \underline{A}_{mL})$$
(8)

defined to be the volumetric integration of the local solid fraction F and is given as between the mushy and liquid cells. The volume-averaged solid fraction $\langle F \rangle_m$ is ated at the face center shown in Figure 1. \underline{U}_{mL} is the convective velocity at the face In the above equations, \underline{A}_{mL} is the outward normal vector from the cell Ω_m and is directed toward the neighboring liquid cell. The associated enthalpy H_{mL} is evalu-

$$\langle F \rangle_m \equiv \frac{1}{\|\Omega_m\|} \int_{\Omega_m} F \, d\Lambda$$
 (9)

It is noted that $\langle \rho \rangle_{\Omega_m}$ is a time-dependent property, since $\langle F \rangle_m$ varies with the time. Thus, Eq. (7) can be written as

$$\sum \rho_L \underline{U}_{mL} \bullet \underline{A}_{mL} = -\|\Omega_m\| \frac{\partial}{\partial t} \langle \rho \rangle_{\Omega_m} \tag{10}$$

the corrected velocities becomes exactly equal to the right-hand side of Eq. (10). then corrected by using a scaling factor such that the net mass flow rate due to averaged velocity magnitudes for the two adjacent cells. Each estimated velocity is mushy and liquid cells. The velocity at this interface is initially estimated to be the It is worth noting that Eq. (10) is the constraint prescribed at the interface of the

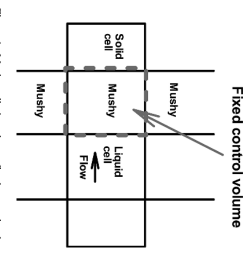


Figure 1. Mushy cell viewed as a fixed control volume

simulation in the liquid region. The interface velocity thus obtained serves as the boundary condition for the flow

condition relating to the enthalpy of solid or liquid is commonly required, To complete the derivation of mushy cell tracking equation, one additional

$$\begin{cases}
H_L \equiv C_s T_m + \underbrace{\Delta H}_{\text{Latent heat}} + C_L (T_L - T_m) & \Omega_L \cup \partial \Omega_{mL} \\
H_s \equiv C_s T_s & \Omega_s \cup \partial \Omega_{ms}
\end{cases}$$
(11)

leads to $H_{mL} = C_s T_m + \Delta H$. For convenience, it can be assumed that $T_{ms} = T_m$. These specifications will provide the essential boundary condition at the interface its temperature becomes the same as T_m . Combining Eqs. (8), (10), and (11), we have of the mushy and liquid (or solid) cells. In other words, once the mushy cell is formed It should be noted that the interface temperature T_{mL} is set as T_m in Eq. (11), which

$$\|\Omega_{m}\| \left(\frac{\partial}{\partial t} \langle \rho H \rangle_{\Omega_{m}} - H_{mL} \frac{\partial}{\partial t} \langle \rho \rangle_{\Omega_{m}} \right)$$

$$= \sum_{m} (k_{s} \langle \underline{\nabla} T_{s} \rangle \bullet \underline{A}_{ms}) + \sum_{m} (k_{L} \langle \underline{\nabla} T_{L} \rangle \bullet \underline{A}_{mL})$$
(12)

The volume-averaged properties $\langle \rho H \rangle_{\Omega_m}$ and $\langle \rho \rangle_{\Omega_m}$ seen above are given as follows:

$$\begin{cases} \langle \rho H \rangle_{\Omega_m} = & \underbrace{\langle F \rangle_m (\rho_s C_s T_m)} + \underbrace{(1 - \langle F \rangle_m) \rho_L (C_s T_m + \Delta H)}_{\text{Enthalpy of solid part}} + \underbrace{(1 - \langle F \rangle_m) \rho_L (C_s T_m + \Delta H)}_{\text{Enthalpy of liquid part}} \end{cases}$$
(13)

Under these conditions, Eq. (12) can be simplified as

$$\begin{split} &\|\Omega_m\| \left\{ \begin{bmatrix} \rho_s C_s T_m \frac{\partial \langle F \rangle_m}{\partial t} - \rho_L (C_s T_m + \Delta H) \frac{\partial \langle F \rangle_m}{\partial t} \end{bmatrix} \right\} \\ &- (C_s T_m + \Delta H) \left(\rho_s \frac{\partial \langle F \rangle_m}{\partial t} - \rho_L \frac{\partial \langle F \rangle_m}{\partial t} \right) \\ &= \sum (K_s \langle \nabla T_s \rangle \bullet \underline{A}_{ms}) + \sum (K_L \langle \nabla T_L \rangle \bullet \underline{A}_{mL}) \end{split}$$

or.

$$-\|\Omega_m\|(\rho_s \Delta H) \frac{\partial \langle F \rangle_m}{\partial t} = \sum (K_s \langle \overline{\nabla} T_s \rangle \bullet \underline{A}_{ms}) + \sum (K_L \langle \overline{\nabla} T_L \rangle \bullet \underline{A}_{mL})$$
 (14)

Further manipulation of Eq. (14) results in

$$\|\Omega_m\|\frac{\partial \langle F\rangle_m}{\partial t} = \sum \left(\frac{-K_s \langle \overline{\nabla} T_s \rangle \bullet \underline{A}_{ms}}{\rho_s \, \Delta H}\right) + \sum \left(\frac{-K_L \langle \overline{\nabla} T_L \rangle \bullet \underline{A}_{mL}}{\rho_s \, \Delta H}\right)$$

01.

$$\|\Omega_m\| \frac{\partial \langle F \rangle_m}{\partial t} = \sum_{f_{i=L,s}} \left(\frac{-K_i}{\rho_s \, \Delta H} \langle \nabla T_i \rangle \bullet \underline{A}_f \right) \tag{15}$$

where \underline{A}_f is the outward normal vector of face_f from the mushy cell Ω_m and is directed

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mentioned assumptions. fluxes from the liquid and solid phases will be included, due to the above two main toward the neighboring cell of i (solid or liquid phase). Only the thermal diffusion

addition, the associated essential boundary conditions are as follows: so that it can be used to define the new front location at the updated time level. In mushy cell. This minimum time step is determined by scrutinizing every mushy cell ing equation is the minimum time step which is applicable to update $\langle F \rangle_m$ for every As stated in Part I of [5, 6], the main reason for choosing the mushy cell track-

$$T_{mL} = T_{ms} = T_m$$

$$\sum \rho_L \underline{V}_{mL} \bullet \underline{A}_{mL} = -\|\Omega_m\| \frac{\partial}{\partial t} \langle \rho \rangle_{\Omega_m}$$
(16)

sby-cell manner as illustrated in Figure 2. In a similar manner, one can derive the the newly solidified mushy cells. Thus, the mushy cells can be advanced in a cellmushy cells can be easily identified by examining the neighbors (in liquid state) of cell-by-cell manner to update $\langle F \rangle_m$ that the advancing length of the moving front is limited by the length of a cell in a where T_{mL} is the temperature between the mushy and liquid cells. It is worth noting between mushy and liquid or solid phases [6]. get the Stefan condition shown in Eq. (4) by essential temperature boundary for every mushy cell. The candidates for the conditions at the the moving-control-volume

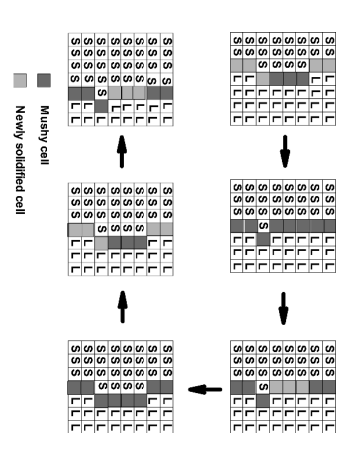


Figure 2. Illustration of the mushy cell advancing.

3. NUMERICAL METHOD

cell boundary. It is also worthwhile to note the following advantages of storing field still retained because all variables are evaluated at the cell center rather than on a tracking algorithm, it is worth noting that the convection terms near the walls are and nodes, respectively. Neighboring cells are defined as those sharing a common the sake of illustration. hedrons or cells. The boundaries and vertices of the cells are referred to as the faces variables at the cell center: Figure 3. The thermally driven mushy cell movement is described in Figure 2 for face. All transport variables are stored at the cell centers. The resulting cells are The domain to be investigated is initially subdivided into several convex polynonstaggered cell-centered control volumes, When applying the cell-by-cell thermally driven mushy cell as illustrated

- moving front. Therefore, the computational time can be considerably reduced. The mushy cell can be easily identified without interpolating or extrapolating the
- Heat flux into the mushy cell can be easily calculated.
- easily dealt with. The number of faces of one particular cell is fixed so that the matrix solver can be
- temperature at the cell center instead of the melting or frozen temperature at the The boundary temperature along the moving front could be easily set as the mushy front. The interpolated or extrapolated procedures can therefore be omitted.

form by choosing the appropriate Γ and S_{Φ} : The transport equation for a scalar quantity Φ can be cast into the following

$$\frac{\partial(\rho\Phi)}{\partial t} + \frac{\partial}{\partial x_i}(\rho u_i\Phi) = \frac{\partial}{\partial x_i}\left(\Gamma\frac{\partial\Phi}{\partial x_i}\right) + S_{\Phi}$$
 (17)

where Γ is the transport coefficient such as the viscosity μ and the ratio $\alpha_L = k_L/C_L$.

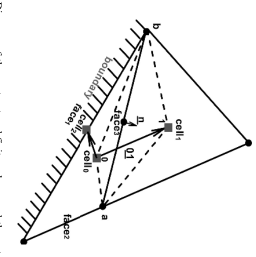


Figure 3. Diagram of the unstructured finite-volume and the notations used.

in the following equation: Integration of Eq. (17) over the control volume Ω_0 , schematic in Figure 3, can result

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$$\frac{\|\Omega_0\|\rho\Phi_0''^{+1}}{\delta_t} + \sum_f J_f \Phi_f = \sum_f D_f + S_{\Phi} \|\Omega_0\| + \frac{\|\Omega_0\|\rho\Phi_0''}{\delta_t}$$
 (18)

order temporal accuracy. the domain Ω_0 . In Eq. (18), the Euler time-stepping method is used to render firstrepresents the diffusion transport through the face "face_f". $\|\Omega_0\|$ is the volume of The mass flow rate J_f is defined to be larger than zero for the fluid leaving Ω_0 . D_f

value of Φ_f at the cell face could be calculated from the following formula: dients can be devised by virtue of the divergence theorem, a hyper-order-accurate dependent on the cell shape. If a shape-independent formulation [23] for the cell graleast-squares technique [24]. These techniques are, however, quite expensive and are eral gradient reconstruction strategies have been proposed [24–26]. Among them, the obtained directly from the Taylor series expansion. To overcome this difficulty, sev-CVFEM method [18] employed the shape function, and another approach used the In the absence of a line structure, the field variable and its derivatives cannot be

$$\Phi_f = \underbrace{\Phi_{\text{upwind}}^{n+1}}_{\text{1st term}} + \underbrace{\nabla \Phi_{\text{upwind, r}}^n \bullet d\underline{r}}_{\text{2nd term}}$$
(19)

cell, and the reconstructed gradient $\overline{\nabla}\Phi_{\text{upwind, r}}^n$ is evaluated from the upwind cell at the time instant t_n . Note that all field variables are treated implicitly, while their gradients are evaluated explicitly. upwind cell to the center of the face. The value of $\Phi_{\rm upwind}^{n+1}$ is obtained from an upwind In the above equation, $d\underline{r}$ denotes the vector with direction from the centroid of an

based on the Gauss divergence theorem: The transport flux D_f due to heat conduction through face_f is derived below

$$D_f \equiv \alpha_i \langle \overline{\Sigma} T \rangle_f \bullet \underline{A}_f \tag{20}$$

flux across a face can be generally expressed as [23] where the thermal diffusivity α_i is defined as $\alpha_i = k_i/C_i$. In Eq. (20), the temperature

$$\underbrace{\langle \overline{\Sigma}T \rangle_{f} \bullet \underline{A}_{f} = (T_{1} - T_{0}) \underbrace{\frac{\underline{A}_{f} \bullet \underline{A}_{f}}{\underline{A}_{f} \bullet 0\underline{1}}}_{\text{first term}} + \underbrace{\langle \overline{\Sigma}T \rangle_{f} \bullet \left(\underline{A}_{f} - \underline{0}\underline{1} \underbrace{\frac{\underline{A}_{f} \bullet \underline{A}_{f}}{\underline{A}_{f} \bullet 0\underline{1}}}\right)}_{\text{Second term}} \tag{21}$$

where

$$\langle \overline{\Sigma}T \rangle_f = rac{1}{2} \left(\langle \underline{\nabla}T \rangle_0 + \langle \underline{\nabla}T \rangle_1
ight) \qquad ext{and} \qquad \langle \underline{\nabla}T \rangle_0 = rac{1}{\|\Omega_0\|} \sum_f T_f \underline{A}_f$$

cells. The face vector \underline{A}_f is directed outward and is normal to face_f for Ω_0 , and $\underline{01}$ is At a face_f, $\langle \nabla T \rangle_f$ and T_f are taken to be the averaged quantities of the two adjacent

able T at the cell centers can be derived in the solid phase or liquid phase under the motion-free condition: Based on Eq. (21), the following linear system of equations for the field vari-

$$\left(\frac{\rho\|\Omega_{0}\|}{\delta t} + B_{0}\right) T_{0}^{n+1} - \sum_{\text{nb}} B_{\text{nb}} T_{\text{nb}}^{n+1} = S_{0} + \frac{\rho\|\Omega_{0}\|}{\delta t} T_{0}^{n}
B_{\text{nb}} = \alpha_{i} \frac{A_{f} \bullet A_{f}}{01 \bullet A_{f}} \qquad B_{0} = \sum_{\text{nb}} B_{\text{nb}}$$
(22)

temperature, respectively. The boundary diffusion flux can then be treated similarly the mushy and liquid and solid cells is specified by the constant melting or freezing center of the boundary. When solving Eq. (22), the interface temperature between contains the super-linear term, such as the secondary diffusion flux terms in Eq. (21). Here, the summation is made over all the neighbors "nb" of Ω_0 . The source term S_0 to that at an interior face using Eq. (21). The flux D_2 on a boundary schematic in The diffusion fluxes at the boundaries are also included in S_0 , and T is applied at the Figure 2 can be written as

$$D_{2} = \alpha_{i}(T_{2} - T_{0}) \underbrace{\frac{A_{f} \bullet A_{f}}{A_{f} \bullet 02}}_{\underbrace{A_{f} \bullet 02}} + \underbrace{\alpha_{i}(\overline{\nabla}T)}_{f} \bullet \left(\underbrace{A_{f} - 02}_{\underbrace{A_{f} \bullet 02}} \underbrace{A_{f} \bullet A_{f}}_{\underbrace{A_{f} \bullet 02}}\right)$$
(23)

is included in the right-hand side of Eq. (22). For the case with Neumann boundary Dirichlet-type boundary condition, the first term in Eq. (23) is treated implicitly and from the center of the cell Ω_0 to the centroid of the boundary face. For the case with where T_2 is evaluated at the center of the boundary face and vector $\underline{02}$ is measured integration. condition, the specified flux can be implemented directly in the control-volume will be included in the left-hand side of Eq. (22). As for the second term in Eq. (23), it

and source terms, the following linear system of equations for Φ can be derived at each cell center in the liquid phase containing the convection effect: After assembling all the discretized terms, including the convection, diffusion,

$$\frac{\|\Omega_0\|\rho\Phi_0^{k+1}}{\delta t} + B_0\Phi_0^{k+1} - \sum_{\text{nb}} B_{\text{nb}}\Phi_{\text{nb}}^{k+1} = S_0 + \frac{\|\Omega_0\|\rho\Phi_0^k}{\delta t}$$
(24)

included in S_0 , and B_0 is the coefficient containing the contributions from the correction of Φ_f , as shown in Eq. (19). The flux contributions at boundaries are cells of Ω_0 . The source term S_0 contains the volumetric source of Φ and the deferred Here, $\|\Omega_0\|$ is the volume of Ω_0 . The notation nb denotes the number of neighboring

The superscript k denotes the k_t^{th} sub-time step in the n time step. convection and diffusion fluxes in the liquid phase considering the convection effect.

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At the face "face," schematic in Figure 1, the mass flow rate is determined by drawbacks, a scheme [17, 23, 27] similar to that of Rhie and Chow [28] will be used prone to oscillatory modes and checkerboard pressure patterns. To overcome these the mass flow rate J_f , which is obtained by averaging the cell center velocities, is Since the pressure and the velocity components are stored at the cell centers,

$$J_{f} = \rho_{f} \underbrace{A_{f}} \bullet \left(\frac{\underline{U}_{0}^{*} + \underline{U}_{1}^{*}}{2}\right)$$

$$-\underbrace{\frac{\rho_{f}(\|\Omega_{0}\| + \|\Omega_{1}\|)}{(\rho\|\Omega_{0}\|/\delta t + B_{0}) + (\rho\|\Omega_{1}\|/\delta t + B_{1})}}_{\text{correction term from momentum}} \left[\left((P_{1} - P_{0}) - \langle \overline{\nabla}P \rangle_{f} \bullet \underline{01}\right) \underbrace{\frac{A_{f} \bullet A_{f}}{A_{f} \bullet \underline{01}}}\right]$$

(25)

coefficient in the discrete momentum equation for the cell Ω_0 shown in Eq. (25), where \underline{U}^* is the velocity field that satisfies the momentum conservation law. B_0 is the $\langle \overline{\nabla} P \rangle_f$ is the averaged pressure gradient evaluated at the cell face "face," and $\underline{01}$ vector measured from the center of Ω_0 to the center of Ω_1 .

the standard diffusion term: the thermal buoyancy force, and the stress tensor components are not included in In each momentum equation, the source term contains the pressure gradient,

$$\frac{\partial}{\partial x_j} \left(\mu \frac{\partial u_j}{\partial x_i} - \frac{2}{3} \delta_{ij} \mu \frac{\partial u_l}{\partial x_l} - \delta_{ij} P \right) - \rho g_i \beta_T (T - T_m)$$
 (26)

Integration of Eq. (26) over the control volume Ω_0 results in the following equation:

$$\sum_{f} \left(\mu_f \frac{\partial u_j}{\partial x_i} - \frac{2}{3} \delta_{ij} \mu_f \frac{\partial u_l}{\partial x_l} - \delta_{ij} P_f \right) A_j + \left[-\rho g_i \beta_T (T - T_m) \right] \| \Omega_0 \|$$
 (27)

yancy force due to thermal expansion is also evaluated at the control-volume center. evaluated by averaging the cell derivatives and cell values, respectively. The buowhere the viscosity μ_f , the extra shear, and the pressure forces at the cell face are

be derived as follows, according to [27]: Within the SIMPLE solution framework, the pressure-correction equation can

$$\underline{U}' = -\frac{\delta t}{\rho} \underline{\nabla} P' \tag{28}$$

$$= P^k + P' \qquad \underline{U}^{k+1} = \underline{U}^k + \underline{U}' \tag{29}$$

$$\underline{\nabla} \bullet (\rho \underline{U}') = -\underline{\nabla} \bullet (\rho \underline{U}^k) \tag{30}$$

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Substituting Eq. (28) into Eq. (30), the pressure-correction equation is derived as

$$\underline{\nabla}^2 P' = \frac{1}{\delta t} \underline{\nabla} \bullet \left(\rho \underline{U}^k \right) \tag{31}$$

where the superscript k represents the predictor step or the previous corrector step. After discretization and integration of Eq. (31) over the control volume Ω_0 , the following discrete equation can be obtained:

$$A_{0}P'_{0} + \sum_{\text{nb}} A_{\text{nb}P'_{\text{nb}}} = -\frac{1}{\delta t} \sum_{j} (\rho \underline{U}^{k} \bullet \underline{A})_{f} + \sum_{f} \left[\langle \overline{\nabla} P' \rangle_{f} \bullet \left(\underline{A}_{f} - \underline{01} \frac{\underline{A}_{f} \bullet \underline{A}_{f}}{\underline{A}_{f} \bullet \underline{01}} \right) \right]$$

$$A_{\text{nb}} = -\frac{\underline{A}_{f} \bullet \underline{A}_{f}}{\underline{01} \bullet \underline{A}_{f}} \qquad A_{0} = -\sum_{\text{nb}} A_{\text{nb}}$$

$$(32)$$

then the Neumann-type boundary condition [8], Eq. (28), is employed along with the corrector step is repeated until convergence. If the boundary pressure is not known, solver, both velocity and pressure fields can be updated using Eq. (29). The entire tion. Once the solution of Eq. (32) is obtained through the preconditioned CG metric and positive definite. The preconditioned BiCGSTAB method is selected the pressure-correction coefficient matrix equation, since it is classified to be symemployed in the present study. The preconditioned CG method is used to solve gradient method. known boundary velocity. The solution procedures based on the liquid phase confor solving the momentum and the discretized temperature equations with convectaining the convection effect for solidification are summarized as follows. The above system of equations will be solved using a preconditioned conjugate Two algorithms of the conjugate gradient (CG) type [19] are

- Define spatially the liquid phase, the mushy zone, and the solid phase. The minimum time step δt is then determined from the mushy cell tracking equation. The
- 2 scale. Thus, the sub-time step $(\delta t)_s$ is chosen as length h, $(\delta t)_{\text{ref}} = 0.3 (h^2 \rho_L C_L / k_L)$ is currently selected as a constraint on $(\delta t)_{s}$, cell-by-cell approach is used to move the front by virtue of the characteristic cell tracking equation, the velocity is controlled by the diffusion contribution. Since a δt is decomposed into several sub-time steps, i.e., $\delta t = \sum_{s} (\delta t)_{s}$. In the mushy cell to determine the velocity, pressure, and temperature. To stabilize the calculation, and the governing equations in the liquid phase using the SIMPLE algorithm value of $\langle F \rangle_m$ can be then updated for each mushy cell. Solve for the heat conduction equation in the solid phase with the time step δt $h^2 \rho_L C_L/k_L$ has the unit of time and is therefore considered as the local cell's time is the sub-time step of the minimum time step δt .

$$\delta t = k(\delta t)_{\text{ref}} + (\delta t)_{k+1} \quad 0 < (\delta t)_{k+1} < (\delta t)_{\text{ref}}$$

$$(\delta t)_{s} = \begin{cases} \delta t & \text{for } \delta t < (\delta t)_{\text{ref}} \\ \frac{\delta t}{k+1} & \text{for } \delta t > (\delta t)_{\text{ref}}, s = 1, 2, \dots, k+1 \end{cases}$$
(33)

For every single sub-time step $(\delta t)_s$:

- (e) equations.
- 6 Substitute the new velocities into the pressure-correction equation and then magnitudes, which satisfy the mass conservation law. solve the pressure-correction equation to update the velocity and pressure
- Use the updated velocity vector to solve for the temperature
- $\overline{\mathcal{S}}$ Go to step (a).

converge to the user specified tolerance. Repeat this procedure until the values of pressure, temperature, and velocity

 $\dot{\mathfrak{S}}$ Go to step 1 for the next time step. If the solid phase cannot move farther into a mushy cell, then stop the solidification process

4. NUMERICAL EXAMPLES

to clarify whether the solution is physically correct. problem will be studied systematically using the present mushy cell tracking method experimental results than with the predicted fine-grid and high-order solution. coarse-grid solution obtained from the lower-order scheme agrees better with the ancy is associated with the myth of grid-converged solution in the sense the numerical result was not consistent with the experimental results. The discrepexperimentally [1] and numerically [2-4, 11, cally [4]. After validating these two benchmark tests, the gallium melting studied chosen to access the present method. effect. The first problem is also known as the analytic Stefan problem [14] and will be ing in a rectangular cavity heated from the side wall with consideration of convection as the 1-D solidification without consideration of the convection effect, and tin melt-Two model problems are used to validate the present method. They are known The second 13, problem has been studied numeri-15, 16] is investigated. However, that the

4.1. Solidification of One-Region Problem without Convection

predicted solid-liquid interface. The domain and the specified boundary conditions An example taken from [14, 21] is investigated for evaluating the quality of the

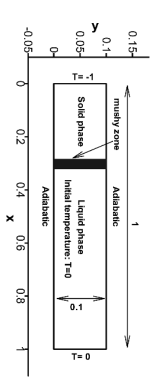


Figure 4. Schematic of the physical domain and the specified boundary conditions for the one-region

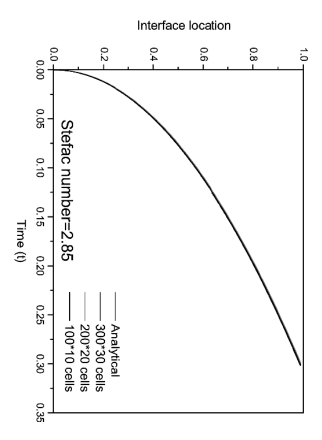


Figure 5. Comparison of the simulated and analytical interface locations for the investigated 1-D, one-region solidification problem (St = 2.85).

are shown in Figure 4. 200×20 , and 300×30 resolutions are selected for the current numerical simulation. Under $\rho_s = \rho_L = 1$, $k_s =$ Three orthogonal and quadrilateral meshes with 100×10 , k_L 1, and $C_s =$ 1, the latent heat ΔH will be used

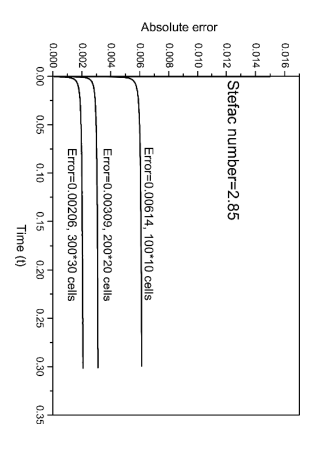


Figure 6. Simulated absolute errors for the numerical and analytical solid-liquid interface locations.

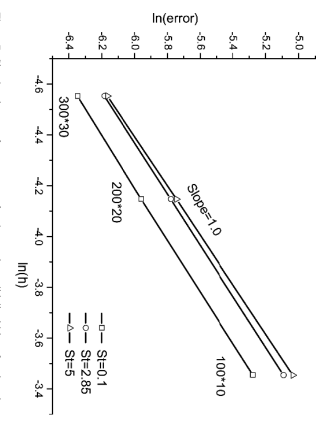


Figure 7. Simulated rates of convergence for the moving solid-liquid interface location

crepancy, defined as $\epsilon \equiv$ Figures 5 and 6. The ac at different Stefan numbers are summarized in Table 1. driven mushy cell tracking method is bounded by $\varepsilon \approx O(h)$. The results obtained words, the error of the predicted interface location using the cell-by-cell thermally that the value of n is very close to 1, which can be also seen in Table 1. In other $\varepsilon \approx C(t)h^n$, where h is defined as the characteristic length of a cell. Figure 7 shows and the analytical solutions are perfectly matched in the sense that the predicted dis-S(t) at the Stefan number 2.85 are shown in Figure 5. Clearly, the simulated results η turn out to be 0.4400, 1.7985, and 2.1194, respectively. The simulated results for were performed at the Stefan numbers of 0.1, 2.85, and 5. The resulting values of moving front is derived as $S(t) = \eta \sqrt{t}$ in the Appendix. tained at the constant temperature $T_{\text{surf}} = -1$. The resulting exact solution for the is kept at the freezing (or melting) temperature Initially, the liquid in the domain under investigation is at rest and its temperature adjust the Stefan number St, which is defined as St = as $\varepsilon \equiv \left| S(t)_{\text{num}} - S(t)_{\text{analytic}} \right|$, is negligibly small as shown in The accuracy order is determined according to the relation $T_f = 0$. The wall at x = 0 is main-The current simulations $C_s(T_f - T_{\rm surf})/\Delta H$.

 Table 1. Absolute errors obtained at different Stefan numbers [14, 21]

	300×30	200×20	100×10	Mesh
$S(t)_{\rm analytic} = 0.4400\sqrt{t}$	0.00175	0.00258	0.00510	$\varepsilon(\mathrm{St}=0.1)$
$S(t)_{\rm analytic} = 1.7985\sqrt{t}$	0.00206	0.00309	0.00614	$\varepsilon(\mathrm{St}=2.85)$
$S(t)_{\rm analytic} = 2.1194\sqrt{t}$	0.0021	0.0032	0.0065	$\varepsilon(\mathrm{St}=5)$

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Table 2. Physical data for tin [4]

Units Solid Liquid Symbol kg/m³ 7,500 7,500 ρ w/m K 60 60 J/kg K 200 200 6×10^{-3} NS/m231.9 റ് $kg/m^3 m/S^2 1/K$ 19.1 $\rho g \beta$ 2.67×10^{-4} 1/K 6×10^4 J/kgH

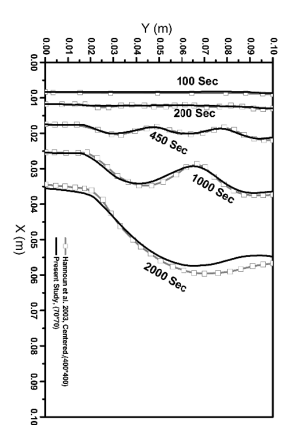


Figure 8. Comparison of the moving fronts during the melting of tin.

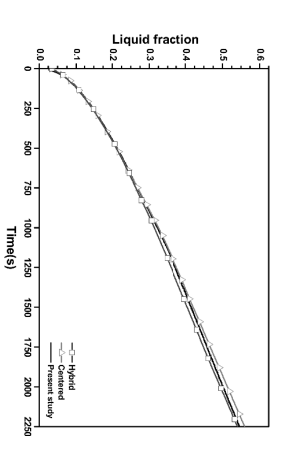


Figure 9. Simulated liquid fractions during the melting of tin (cells: 70×70).

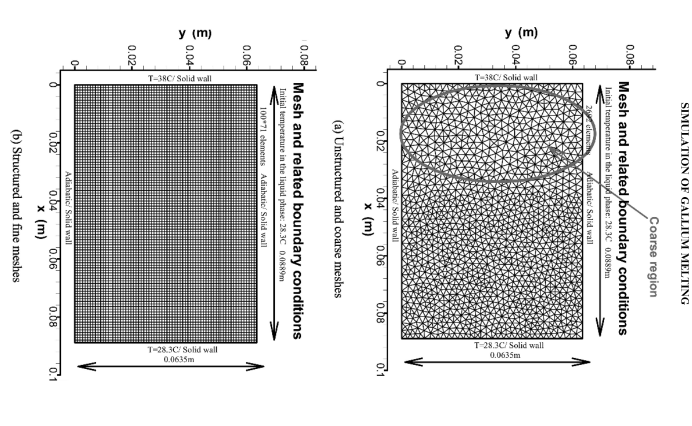


Figure 10. Boundary conditions and computational meshes for simulating gallium melting: (a) unstructured/coarse meshes; (b) structured/fine meshes.

Table 3
Physical
properties
of gallium
[4, 20]

× 10 ⁷	$1.173 \times 10^{-7} 8 \times 10^{-7}$	7.0	29.78	1.8×10^{-3}	395.15	33.5	6,095	Liquid
104	10-4	1	1	0	395.15	33.5	6,095	Solid
J/kg	1/K	$kg/m^3 m/S^2 1/K$	°C	NS/m	J/kgK	$\mathrm{w/m}\mathrm{K}$	kg/m^3	Units
Н	β	$\rho g \beta$	T_m	μ	C_p	k	ρ	Symbol
				,				

4.2. Melting of Tin

505 K, respectively, and the bottom and top boundaries are assumed to be thermally cal data [4] listed in Table 2. The left and right wall temperatures are 508 K and with those given in [4], as shown in Figure 8 and Figure 9. results regarding the moving front location and the liquid fraction compare all well The melting of tin in a square cavity will be investigated at the published physi-The melting temperature of tin is assumed to be 505 K. The computed

4.3. Melting of Gallium

set at T = 28.3°C. are sketched in Figure 10. Initially, the temperature in the solid gallium block is present numerical model. experimental results of Gau and Viskanta [1] is chosen for the verification of the studied experimentally [1] and numerically [2value of $T=28.3^{\circ}\mathrm{C}$. The physical properties used in the current calculations are tabulated in Table 3 [4, 20]. $T=38.0^{\circ}\text{C}$, while the right wall temperature remains unchanged with the initial Gallium melting in a rectangular cavity heated from the side wall has been The temperature at the The geometric configuration and boundary conditions left wall is 4, 11, 13, 15, 16]. The widely cited increased instantly to

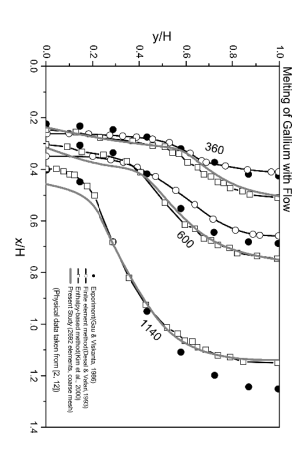


Figure 11. Comparison of the moving interfaces of gallium melting at various time instances.

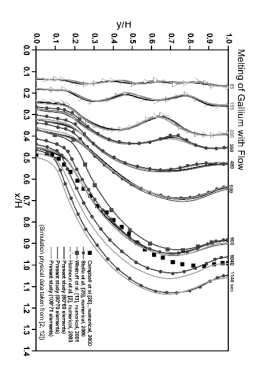


Figure 12. Comparison of the present result with other numerical results

meshes generated around the interface region are very important, since they will tured and structured meshes as depicted in Figure 10 are employed. The coarse shed some light on this controversial prediction, several grid resolutions of unstrucwith the experimental data than those obtained from the fine meshes. In order to predicated dynamics of interface and thermofluid in the coarse meshes match better tization schemes and grid resolutions, to study this problem. Hannoun et al. [4] employed the enthalpy method, together with various discre-They found that the

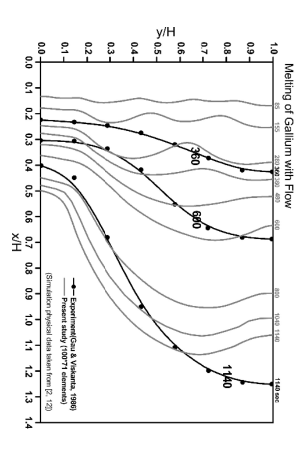


Figure 13. Comparison of the present result with the experimental data [1].

quality compared with the experimental data [1] and other numerical results [2, 3, solutions in the coarse meshes near the interface region can yield better prediction affect the simulated results, as noted by Hannoun et al. [4]. The resulting predicated 11] shown in Figure 11.

well with the numerical results of Hannoun et al. [4] in the early period of simulation numerical results is shown in Figure 12. The currently predicted interface matches $(t < 280 \,\mathrm{s})$ and agrees well with the other numerical results for $t > 360 \,\mathrm{s}$, as shown in lateral meshes are used. Comparison of the currently predicted interface with other flow field and interface dynamics, 80×60 , 90×70 , and 100×71 orthogonal, quadri-To further address the effect of mesh resolution on the predicted thermofluid

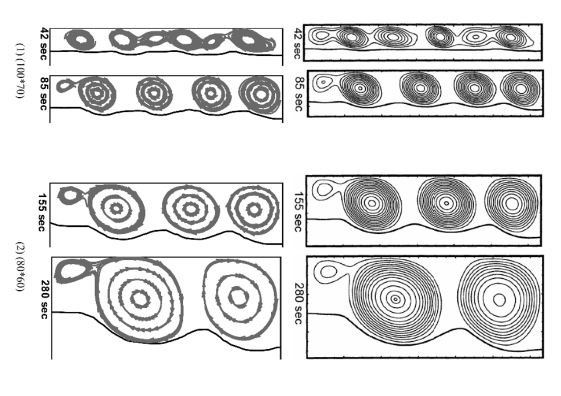


Figure 14. Simulated streamlines: upper row from [4] and lower row from the present study.

time instants are shown in Figure 14. numerical simulation. The corresponding streamline patterns obtained at various discrepancy discussed by Hannoun et al. [4]. This predicted difference might be attributed to the obtained from the fine meshes and the experimental data is significant and has been depicted in Figure 13. Figure 12. Comparison of the numerical results with the experimental data [1] is between the experimental setup and the assumptions made in the The discrepancy between the numerical result and those

5. CONCLUSIONS

the following conclusions can be drawn. convection have been studied to validate the method. Based on the computed results solidification without convection transport, and tin and gallium 2-D melting with so as to be able to accurately predict the movement of the mushy cells. Both 1-D cell tracking equation has been derived to satisfy the mass and energy conservations and liquid-phase regions, with the mushy cells being placed in between. The mushy within the finite-volume context. The physical domain is separated into solid-phase A thermally driven mushy cell tracking solution algorithm has been developed

- moving interface is linear in the sense that $\varepsilon(t) \approx C(t)h$. This finding is close to the order of accuracy predicted by the moving-particle method [12]. agrees well with the analytic solution. The predication accuracy order for the Simulation of the solidification shows that the predicted moving interface S(t)
- 2 assumptions made in the numerical simulation, as elaborated in [4]. might be attributed to the experimental setup, which is not consistent with the tion (t > 360 s). The discrepancy between the experimental and numerical results period ($t = 0-280 \,\mathrm{s}$). The simulated results are different from the experimental mental data [1] and other numerical results [4, 11, 15, 16] in the early simulation thermofluid flow field and dynamics of mushy cells agree well with the experimesh-dependent, but the mesh resolution is not fine enough to resolve the small data [1] and other numerical results [2, 3, 11]. We believe these predictions are the coarse meshes, near the interface region matches better with the experimental As Hannoun et al. [4] demonstrated, our gallium melting interface, predicted in and complex thermofluid flow field structure near the top wall close to the interdata but are consistent with other numerical results in the later period of simulaface area. Simulation of gallium melting in fine meshes shows that the predicted

REFERENCES

- C. Gau and R. Viskanta, Melting and Solidification of a Pure Metal on a Vertical Wall
- Trans. ASME, J. Heat Transfer, vol. 108, pp. 174–181, 1986.

 C. P. Desai and K. Vafai, A Unified Examination of the Melting Process within a Trans-Dimensional Rectangular Cavity, Trans. ASME, J. Heat Transfer, vol. 115, pp. 1072-1075, 1993
- R. Viswanath and Y. Jaluria, A Comparison of Different Solution Methodologies for Melting and Solidification Problems, Numer. Heat Transfer B, vol. 24, pp. 77–105, 1993

372

- 4. Gallium Melting in a Rectangular Cavity Heated from the Side, Numer. Heat Transfer N. Hannoun, V. Alexiades, and T. B, vol. 44, pp. 253–276, 2003. Z. Mai, Resolving the Controversy over Tin and
- S Change Problems—Part I: Concept for Tracking the Moving Front, J. Comput. Mech., Cell-by-Cell Thermally Driven Mushy Cell Tracking Algorithm for Phase-
- 6 in press, 2006. Change Problems—Part II: Phase-Change with Natural Convection, J. Comput. Mech., J. Jan, A Cell-by-Cell Thermally Driven Mushy Cell Tracking Algorithm for Phase-
- 7 W. Shyy, H. S. Udaykumar, M. M. Rao, and R. W. Smith, Computational Fluid Dynamics with Moving Boundaries, Taylor & Francis, Washington, DC, 1996.
- W. Shyy, S. S. Thakur, H. Ouyang, J. Liu, and E. Blosch, Computational Techniques for Complex Transport Phenomena, Cambridge University Press, Cambridge, UK, 1997.
- 9 J. Crank, Free and Moving Boundary Problems, Oxford University Press, Oxford, UK, 1984.
- A. D. Brent, V. R. Voller, and K. J. Reid, Enthalpy-Porosity Technique for Modeling Heat Transfer, vol. 13, pp. 297-318, 1988. Convection-Diffusion Phase-Change: Application to the Melting of a Pure Metal, Numer.
- S. Kim, S. Anghaie, and G. Chen, A Fixed-Grid Two-Phase Numerical Model for Coning for Industrial Multiphase Flows, pp. 27-29, Institut d'Etudes Scientifiques de Cargèse, vection-Dominated Melting and Solidification, Trends in Numerical and Physical Model-
- 12. the Sharp Interface Limit on the Fixed Grids, J. Comput. Phys., vol. 153, pp. 535-574, H. S. Udaykumar, R. Mittal, and W. Shyy, Computation of Solid-Liquid Phase Fronts in
- 13. I. Wintruff, C. Giinther, and A. G. Class, An Interface-Tracking Control-Volume Finite-Application, Numer. Heat Transfer B, vol. 39, pp. 127-149, 2001. Element Method for Melting and Solidification Problems-—Part II: Verification and
- 14. M. N. Ozisik, Heat Conduction, Wiley, New York, 1980.
- 15. F. Stella and M. Giangi, Melting of a Pure Metal on a Vertical Wall: Numerical Simulation, Numer. Heat Transfer A, vol. 38, pp. 193-208, 2000.
- T. A. Campbell and J. N. Koster, Visualization of Liquid/Solid Interface Morphology in Gallium Subject to Natural Convection, J. Crystal Growth, vol. 140, pp. 414–425, 1994.
- 17. J. H. Ferziger and M. Peric, Computational Methods for Fluid Dynamics, Springer-Verlag,
- B. R. Baliga and S. V. Patankar, A Control-Volume Finite Element Method for Two-Dimensional Fluid Flow and Heat Transfer, Numer. Heat Transfer, vol. 6, pp. 245-261,
- 19. Y. Saad, Iterative Methods for Sparse Linear Systems, PWS Publishing, Boston, 1996.
- 20. R. Sampath and N. Zabaras, An Object-Oriented Implementation of a Front Tracking Eng., vol. 44, pp. 1227-1265, 1999. Finite Element Method for Directional Solidification Processes, Int. J. Numer. Meth.
- Solidification Problems—Part I: Method and Validation, Numer. Heat Transfer B, vol. 43, pp. 117–141, 2003. Y. Li, S. V. Garimella, and J. F. Simpson, Fixed-Grid Front-Tracking Algorithm for
- 22. Numer. Heat Transfer, B, accepted, 2007. Y. J. Jan, Thermally Driven Mushy Cell Tracking Algorithm for Dendritic Solidification
- 23 S. R. Mathur and J. Y. Murthy, A Pressure-Based Method for Unstructured Meshes Heat Transfer B, vol. 31, pp. 195-215, 1997
- P. A. Jayantha and I. W. Turner, A Comparison of Gradient Approximations for Use in Finite-Volume Computational Heat Transfer B, vol. 40, pp. 367-390, 2001. Models for Two-Dimensional Diffusion

- 25. G. Labonia, V. Timchenko, J. E. Simpson, S. V. Garimella, E. Leonardi, and G. De Vahl Davis, Reconstruction and advection of a moving interface in three dimensions on a fixed grid, Numer. Heat Transfer B, vol. 34, pp. 121-138, 1998.
- 26. P. Jawahar and H. Kamath, A High-Resolution Procedure for Euler and Navier-Stokes Computations on Unstructured Grids, J. Comput. Phys., vol. 164, pp. 165–203, 2000.
- 27. lating Laminar Reacting Flows, *Numer. Heat Transfer B*, vol. 41, pp. 53–72, 2002. C. M. Rhie and W. L. Chow, Numerical Study of the Turbulent Flow Past an Airfoil with S. Kang and Y. Kim, Pressure Based Unstructured Grid Finite Volume Method for Simu-
- 28 Training Edge Separation, AIAA J., vol. 21, pp. 1525–1532, 1983.

APPENDIX: EVALUATION OF THE MOVING-FRONT LOCATION

Consider the following one-dimensional equation:

$$\frac{\partial T}{\partial t} = \frac{k_s}{\rho_s C_s} \frac{\partial^2 T}{\partial x^2} \qquad 0 < x < H$$

$$\begin{cases}
t = 0, \quad T = T_m \\
x = 0, \quad T = T_{\text{surf}}
\end{cases}$$
(A1)

cular solution is given as where S(t) is the interface between the liquid and solid phase. In Eq. (A1), the parti-

$$T - T_m = C_1 \left[\operatorname{erf}(a) - \operatorname{erf}\left(\frac{x}{\sqrt{4\alpha_s t}}\right) \right]$$
 (A2)

 C_1 and a, could be specified by substituting the boundary conditions into Eq. (A2) to render $T_{\rm surf} - T_m = C_1 \, {\rm erf}(a)$ and $a = S(t)/\sqrt{4\alpha_s}t$. It should be noted that $dT \, [\equiv (\partial T/\partial t) \, dt + (\partial T/\partial x) \, dx]$ at the moving front is zero for the constant-melting or frozen-temperature case. Thus, is defined as $\alpha_s \equiv k_s/\rho_s C_s$. The subscript s represents the solid phase. Two constants. Note that $\operatorname{erf}(a)$ is defined as $\operatorname{erf}(a) = 2 \int_0^a (e^{-\eta^2}) d\eta / \sqrt{\pi}$ and the thermal diffusivity

$$\left(\frac{\partial T}{\partial t}\right)_{x=S(t)} = -\frac{dx}{dt} \left[\left(\frac{\partial T}{\partial x}\right) \right]_{x=S(t)}$$

results. Thanks to the energy conservation principle, we can get

$$\left(\frac{dx}{dt}\right)_{x=S(t)} = \frac{\alpha_s C_s}{(\Delta H)} \left(\frac{\partial T}{\partial x}\right)_{x=S(t)}$$

interface can be described as and C_s is the heat capacity for the solid. Consequently, the equation for the moving in the one-dimensional domain. Note that ΔH is the latent heat of the phase change

$$\left(\frac{\partial T}{\partial t}\right)_{x=S(t)} = -\frac{\alpha_s C_s}{(\Delta H)} \left[\left(\frac{\partial T}{\partial x}\right)^2 \right]_{x=S(t)}$$
(A3)

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Since $(\partial T/\partial t)_{x=S(t)}$ and $(\partial T/\partial x)_{x=S(t)}$ can be respectively expressed as

$$\left(\frac{\partial T}{\partial t}\right)_{x=S(t)} = \frac{C_1 S(t)}{\sqrt{4\alpha_s \pi t^3}} e^{-S(t)^2/4\alpha_s t} \tag{A4}$$

$$\left(\frac{\partial T}{\partial x}\right)_{x=S(t)} = \frac{C_1}{\sqrt{\alpha_s \pi t}} e^{-S(t)^2/4\alpha_s t}$$
(A5)

$$C_1 = -\frac{(\Delta H)a\sqrt{\pi}}{C_s} \left(e^{a^2}\right) \tag{A6}$$

By substituting Eq. (A6) into the equation given by $T_{\rm surf} - T_m = C_1 \, {\rm erf}(a)$, we are led to derive $T_{\rm surf} - T_m = -(\Delta H) a \sqrt{\pi} \left(e^{a^2}\right) \, {\rm erf}(a)/C_s$. To further illustrate the time-varying front feature, the Taylor series approximation is applied to $ae^{a^2} \, {\rm erf}(a)$, thus resulting in

$$ae^{a^{2}}\operatorname{erf}(a) = \frac{2a^{2}}{\sqrt{\pi}} \left(1 + a^{2} + \frac{a^{4}}{2} + \cdots \right) \left(1 - \frac{a^{2}}{3 \times 1!} + \frac{a^{4}}{5 \times 2!} - \cdots \right)$$
$$= \frac{2a^{2}}{\sqrt{\pi}} \left(1 + \frac{2}{3}a^{2} + \frac{4}{15}a^{4} + \cdots \right)$$
(A7)

A further simplification of $T_{\text{surf}} - T_m$ can render

$$T_{\text{surf}} - T_m = -\frac{2(\Delta H)a^2}{C_s} \left(1 + \frac{2}{3}a^2 + \frac{4}{15}a^4 + \cdots\right)$$
 (A8)

signify the importance of the sensible heat relative to the latent heat. If we keep the term $(a)^2 2/3$, the accurate front profile S(t) can be approximated as below after The Stefan number is defined as St $\equiv -(T_{\text{surf}} - T_m)C_s/\Delta H$ and its value is obtained from St $= 2a^2(1 + 2a^2/3 + \cdots)$ for solidification. The Stefan number is defined to some simple calculations.

$$S = \left[\sqrt{-3 + \sqrt{9 + 12 \left(\frac{(T_m - T_{\text{surf}})C_s}{\Delta H} \right)}} \right] \sqrt{\alpha_s t}$$

$$= \left[\sqrt{-3 + \sqrt{9 + 12(St)}} \right] \sqrt{\alpha_s t}$$

$$= \eta \sqrt{t}$$
(A9)

If the simulation is performed at St = 0.1, the values of η becomes 0.4400.