An Improved Algorithm for the Modeling of Vapor Flow in Heat Pipes

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INTRODUCTION

The NASA Lewis Research Center has developed a steady state heat pipe code for use in heat pipe design and analysis of laboratory data, suitable for microcomputers as well as larger machines.\(^1\) It has become apparent that the vapor flow algorithms in this and similar codes\(^2,3\) are not very satisfactory at conditions where compressibility is important. Heat pipes having axially varying heat input, sequential evaporators, or operating near the sonic limit, are not handled well in these codes. Typical of the problems in some codes is the prediction of a negative absolute pressure within an evaporator with large heat input, because of improper treatment of compressibility effects. The ability to handle compressibility and variable heat input is desired for the NASA Lewis code. Therefore, an accurate vapor flow algorithm concise enough to operate on a microcomputer at a reasonable speed and incorporating these features was sought. Finite difference techniques which have been used effectively in fundamental studies of heat pipe vapor dynamics\(^4,5\) were precluded by code size. Among sources consulted in an attack on this problem was the book by Ivanovskii et al.\(^6\) which reviews several studies leading to one-dimensional algorithms for compressible vapor flow in heat pipes.

For the present work, Busse's\(^7\) paper on pressure drop in the vapor phase was used as a starting point. His equations for incompressible flow were readily rederived to include change in density. In addition, the usual assumption of a saturation temperature across each section of the pipe, determined by the local static pressure, was eliminated. In view of the studies of DeMichele,\(^4\) a mean vapor temperature differing from the wall temperature was found from a boundary-layer-type energy equation. The resulting equations were derived so as to be suitable for liquid metal working fluids having two vapor species as well as for fluids with a single vapor species. The algorithm is intended only for the evaporator and adiabatic sections of the heat pipe, because vapor flow in the condenser is complicated by the onset of turbulence and reverse flow.
ANALYSIS

In the analysis to follow, the appropriate features of Busse's method will be introduced, and the development will proceed from there. In addition, some major assumptions for the heat pipe sections being considered should be indicated at the outset: Flow is laminar. The ideal gas law applies to components of the vapor. The equations allow for local chemical equilibrium among species in the vapor. Phase equilibrium occurs only at the wall, so liquid droplets are absent in the vapor. Following Busse, no tangential velocity component exists at the wall in mass addition or removal, the radial pressure gradient may be neglected, and the velocity components are zero at the closed ends of the heat pipe. Also, his method of imposing a negligible radial pressure gradient by equating the mean axial pressure gradient over the cross section to the axial pressure gradient along the center line was used.

Busse began with the axial axisymmetric Navier-Stokes equation

\[ \frac{\partial p}{\partial z} = -\rho \left( u \frac{\partial v}{\partial r} + v \frac{\partial u}{\partial z} \right) - \frac{n \rho}{r} \left( \frac{\partial u}{\partial z} - \frac{\partial v}{\partial r} \right) \]  

where \( p \) is pressure, \( r \) is radial distance, \( u \) is radial velocity, \( v \) is axial velocity, \( z \) is axial distance, \( n \) is dynamic viscosity, and \( \rho \) is density. Busse integrated the mass conservation equation with respect to \( u \) so that this term can be eliminated. This is also done in the present analysis except that density is variable in \( z \) and \( r \):

\[ u = -\frac{1}{r \rho} \int_0^r \frac{\partial}{\partial z} \left( \rho v \right) r' \, dr' \]  

When equation (2) is combined with equation (1) the following results:

\[ \frac{\partial p}{\partial z} = \frac{1}{r} \left[ \frac{\partial v}{\partial r} \left( \int_0^r \rho \frac{\partial v}{\partial z} r' \, dr' + \int_0^r v \frac{\partial u}{\partial z} r' \, dr' \right) \right] - \rho v \frac{\partial v}{\partial z} + \frac{n \rho}{r} \frac{\partial u}{\partial r} \frac{\partial v}{\partial r} \]  

Here, the axial variation of \( u \) has been neglected in the viscous term. The assumption is made that the radial variation of density is much less than the radial variation of the axial velocity since the latter vanishes at the wall. On the basis of this assumption, a presently undefined mean density \( \rho_m \) and its axial derivative are extracted from the integrals, yielding

\[ \frac{\partial \rho}{\partial z} = \frac{1}{r} \left[ \frac{\partial v}{\partial r} \left( \rho_m \int_0^r \rho \frac{\partial u}{\partial z} r' \, dr' + \frac{3 \rho_m}{2} \int_0^r v \frac{\partial r'}{\partial z} r' \, dr' \right) \right] - \rho v \frac{\partial v}{\partial z} + \frac{n \rho}{r} \frac{\partial u}{\partial r} \frac{\partial v}{\partial r} \]  

The crux of Busse's attack on the incompressible case is the assumption of an approximate velocity profile which satisfies all the boundary conditions. It incorporates a parameter \( A(z) \) which may be constant or may vary with position, depending on mass influx boundary conditions. The same profile will be employed in this analysis:

\[ v = 2v_m \left( 1 - \frac{r^2}{R^2} \right) \left[ 1 + A(z) \left( \frac{r^2}{R^2} - \frac{1}{3} \right) \right] \]
Here, $v_m$ is the mean axial velocity and $R$ is the vapor space diameter. When $A(z) = 0$, equation (5) reduces to the Poiseuille profile. Integration of equation (4) over the cross section of the pipe and manipulation gives

$$\frac{dp_m}{dz} = -\frac{4}{3} \frac{d}{dz} \left[ \left( 1 - \frac{A}{6} + \frac{2A^2}{45} \right) \left( v_m^2 p_m \right) \right] - \frac{8n v_m}{R^2} \left( 1 + \frac{2A}{3} \right)$$

(6)

The mean density is assumed to be related to the pressure and a mean temperature $T_m$ by the ideal gas law, which yields upon differentiation

$$\frac{dp_m}{dz} = (GRP) \frac{d}{dz} \left( \ln p_m \right) - (GRP) \frac{d}{dz} \left( \ln T_m \right)$$

(7)

where

$$GRP = 1 - \frac{\partial \ln R}{\partial \ln p}$$

and

$$GRT = 1 + \frac{\partial \ln R}{\partial \ln T_m}$$

and where $R_g$, the gas constant, may depend on pressure and temperature if chemical reactions are considered.

Continuity requires that the vapor mass flux $m(z)$ at any cross section equals the product of mean density $p_m$, the vapor cross-sectional area $A_v$, and the mean axial velocity $v_m(z)$. From this relationship and equation (7) the axial derivative of $v_m$ is found to be:

$$\frac{d}{dz} \left( \frac{v_m}{p_m} \right) = \frac{d}{dz} \left( \frac{m}{p_m A_v} \right) - \frac{d}{dz} \left( \frac{A_v}{p_m} \right) - (GRP) \frac{d}{dz} \left( \ln m \right) + (GRT) \frac{d}{dz} \left( \ln T_m \right)$$

(8)

Equations (6) to (8) are combined and the indicated integrations are performed. Dropping the distinction between the mean cross section pressure and the local pressure by Busse's approximation gives

$$\left[ 1 - \frac{4p_m v_m^2}{3p} \right] \frac{d}{dz} \left( \ln p \right) + \left[ \frac{4p_m v_m^2}{3p} \right] \frac{d}{dz} \left( \ln m \right) - \frac{4p_m v_m^2}{3p} \left( GB \right) \frac{d}{dz} \left( A_v \right)$$

(9)

where

$$GB = \left( 1 - \frac{A}{6} + \frac{2A^2}{45} \right)$$

and

$$HB = -\left( 1 - \frac{8A/15}{6} \right)$$

The variation of $A_v$ with $z$ is understood to be small.

The mean temperature $T_m$ is evaluated by means of an energy integral. Figure 1 shows the control volume through which energy is transported. The energy flows into the boundary are equated with the energy flows out, internal energy and pressure are eliminated by using enthalpy, and the limit $\Delta z \to 0$ is taken, to give
Figure 1. Control volume for the integral energy equation.

\[ \int_0^R \left[ \frac{d}{dz} \rho v \left( h + \frac{v^2}{2} \right) \right] r \ dr = \int_0^R \theta r \ dr + q_R R - p R U_R \]  \hspace{1cm} (10)

where \( h \) is the enthalpy of the mixture per unit mass, \( q_R \) is the energy flux entering from the wall of the pipe, and \( \theta \) is the viscous dissipation per unit volume. The relation for \( u_R \), the wall radial velocity, is given by the mass conservation equation in the form

\[ u_R = -\frac{1}{R \rho R} \int_0^R \frac{d}{dz} (\rho v) r \ dr \]  \hspace{1cm} (11)

The energy influx \( q_R \) is specified as \(-p R U_R (e_R + \frac{u_R^2}{2})\), where \( e_R \) is internal energy. The result of combining equations (10) and (11) is

\[ \int_0^R \left( h + \frac{v^2}{2} - H_R \right) \frac{d}{dz} (\rho v) r \ dr + \int_0^R \rho v \frac{d h}{dz} r \ dr + \int_0^R \rho v^2 \frac{d v}{dz} r \ dr = \int_0^R \theta r \ dr \]  \hspace{1cm} (12)

The mean value of both the enthalpy and the density can be separately extracted from the integrals if the assumption is again made that the variation of both variables with radius is much less than that of axial velocity. When these mean values have been extracted, and terms common to all expressions cancelled, the energy equation becomes

\[ \left( h_m - H_R \right) \frac{d}{dz} \rho m v m + \frac{1}{R^2} \left( \int_0^R v^3 r \ dr \right) \frac{d \rho m}{dz} + \rho m v m \frac{d h_m}{dz} + \rho m \int_0^R \frac{d v^3}{dz} r \ dr = \frac{2}{R^2} \int_0^R \theta r \ dr \]  \hspace{1cm} (13)

where

\[ H_R = h_R + \frac{u_R^2}{2} \]
If chemical equilibrium in the vapor must be considered, enthalpy $h_m$ becomes a function of both $T_m$ and $p$. When its derivative is inserted into equation (13) with equations (7) and (9), and the integrations are performed, the result is

$$
\left[ \frac{3h_m}{3 \ln p} \right]_{T_m} - F_3(\text{GRP}) \frac{d \ln p}{dz} + \left[ \frac{3h_m}{3 \ln T_m} \right]_p + F_3(\text{GRT}) \frac{d \ln T_m}{dz} + (\text{HAF}) \frac{dA}{dz}
$$

$$
= 8n \left[ \frac{(1 + 2A^2/9)v_m}{\rho R^2} \right] - (\text{HVE}) \left( \frac{d \ln m}{dz} - \frac{d \ln A}{dz} \right)
$$

where

$$F_3 = 8 \left( \frac{1}{4} - \frac{A}{10} + \frac{A^2}{30} - \frac{2A^3}{945} \right)$$

$$\text{HAF} = -\frac{2}{5} \left( 1 - \frac{2A}{3} + \frac{4A^2}{53} \right)$$

$$\text{HVE} = h_m - H_R + \left( \frac{3F_3 v_m^3}{2} \right)$$

An additional equation required to find the three dependent variables $p$, $A$, and $T_m$ comes from Busse's approximation that the axial gradient of the mean pressure equals the center line pressure gradient. The latter, from equation (1) for $r = 0$ using equations (4) and (5), is

$$
1 - \frac{4\rho v_m^2}{p} (1 - A/3)^2(\text{GRP}) \frac{d \ln p}{dz} + \left[ \frac{4\rho v_m^2}{p} (1 - A/3)^2(\text{GRT}) \right] \frac{d \ln T_m}{dz}
$$

$$
- \left[ \frac{4\rho v_m^2}{3p} (1 - A/3) \right] \frac{dA}{dz} = -\frac{8n v_m}{\rho R^2} (1 - 4A/3)
$$

$$
- \left[ \frac{4\rho v_m^2}{p} (1 - A/3)^2 \right] \left( \frac{d \ln m}{dz} - \frac{d \ln A}{dz} \right)
$$

The mass flux is related to the heat flux into the pipe by

$$
dq_R = \frac{1}{2\pi R} \frac{dm}{dz} \left[ h_{v_f} + \left( \frac{1}{2\pi R R} \frac{dm}{dz} \right)^2 \right]
$$

where the kinetic energy of the injected vapor has been incorporated, and $h_{v_f}$ is the heat of evaporation. Equations (9), (14), and (15), with equation (16) and appropriate boundary conditions enable a solution for $p$, $A$, and $T_m$. 

5
The equations derived here pertain to alkali metal working fluids if equilibrium between monomeric and dimeric species is experienced. With proper application, they also apply to working fluids of a single gaseous species. If, in the case of the alkali metals, composition remains fixed from the point of injection, the equations must be modified. This case has been considered but the results are much less satisfactory than for equilibrium and are not presented here.

RESULTS AND DISCUSSION

To illustrate the application of the algorithm, several cases were run involving alkali liquid metal working fluids. Entropies and enthalpies of the species referred to a common base that includes chemical energy were used with polynomials for the specific heats to find thermodynamic properties. These data enabled the determination of vapor pressure, heat of vaporization, the first derivatives of composition and composition of the vapor phase. The latter includes both monomers (atoms) and dimers (molecules) in the case of alkali metals. The assumption is made that the ideal gas law applies to each of the vapor species. While it would be desirable to incorporate real gas effects in the form of virial equations, this would greatly complicate the computations.

The three equations involving the first derivatives of \( p \), \( A \), and \( T_m \) were solved by a simple Runge-Kutta routine for the set of simultaneous equations (9), (14), and (15). No study of the effect of step size on the precision of results was made. Each computation began with 20 uniform steps in the evaporator, but step size was reduced as the slope of \( p \) became increasingly steep in the vicinity of the sonic limit.

Figure 2 illustrates a simple application of the algorithm which solves the equation set. The heat input to a sodium-filled heat pipe of 0.02 m diameter with an evaporator length of 0.1 m was increased until the pipe was choked at an evaporator endcap vapor temperature of 900 K and a heat input of 5870 W. The sonic limit computed from the approximation of Busse was about 5580 W. Figure 2 shows the profiles of vapor mean temperature \( T_m \), wall temperature \( T_w \), the velocity profile parameter \( A \), and the pressure ratio \( p/p_0 \) through the evaporator, where subscript 0 denotes the vapor condition at \( z = 0 \), the endcap. Figure 3 shows the velocity profile \( v/v_m \) at the downstream end of the evaporator where choking occurs, with a Poiseuille profile for comparison. The maximum value did not occur at the centerline, as expected. Busse's approximate profile is a three-point fit, so the profile shown in Figure 3 must be regarded as the best representation of a very flat profile available under the circumstances.

In the next example two evaporator sections in sequence were considered, the first 0.2 m long, the second 0.1 m long, separated by an adiabatic section of 0.2 m length. The working fluid was sodium and the evaporator end cap temperature was 1000 K. Both evaporator sections were receiving heat at the rate of 71 600 W/m, which resulted in choke at the end of the second evaporator. Figure 4 shows the profiles of temperature, pressure ratio, and \( A \). The scale of the figure is too small to show clearly the decline in the parameter \( A \) in the adiabatic section from 0.738 to 0.704. In incompressible flow, the limit for \( A \) in very long adiabatic sections should decline toward zero, the Poiseuille condition.
Figure 2. Variation of heat pipe parameters with location.

Figure 3. Comparison of \( V/V_m \) for Poiseuille and choked flow conditions.

Figure 4. Variation of heat pipe parameters with location for a multi-location heat source on evaporator.
Another example shows the sonic limit heat transport against evaporator endcap temperature for a sodium-filled evaporator of 0.02 m diameter and 0.1 m length. The short evaporator length minimized the viscous pressure loss, so the results can be compared to those computed from inviscid equations of Busse and Levy. A simple control structure appended to the vapor pressure subroutine generated figure 5 in a short time.

The most suitable data against which to check the algorithm are those of Ivanovskii et al. They inserted a temperature microprobe into a sodium heat pipe having an evaporator length of 0.1 m, an adiabatic section length of 0.05 m, a condenser length of 0.55 m, and a diameter of 0.014 m. Temperatures were read at intervals along the axis with the probe. Liquid-saturated wicking on the probe was to ensure that the temperatures being observed were the same as those at the vapor-liquid interface. No details were given concerning any effect of the pressure field around the probe tip on the measured temperature. For an input of about 500 W, stated to be accurate to within 5 to 10 percent, the data are shown in figure 6. Also shown are lines of $T_m$ and $T_R$ computed by the algorithm. Equations (9), (14), and (15) were used only in the evaporator and adiabatic sections. An extension into the condenser was made by equating $T_m$ to $T_R$ beyond the point where $T_m$ exceeded $T_R$. The justification is the observation that at some point in the condenser, $T_m$ must converge to $T_R$ as it did at the beginning of the evaporator. For this portion of the solution, the equations were suitably modified to yield as dependent variables only $p$ and $A$. This simple procedure, enabling the solution to be continued into the condenser, neglected the known complications of condensation shocks and recirculation.

For the same heat pipe at a heat input of 1000 W, figure 7 shows experimental data of Ivanovskii and lines of $T_m$ and $T_R$ from the algorithm. Also shown is a curve of $T_R$ computed by Chen and Faghri using the Phoenix code. It is in basic agreement with the results computed by the algorithm presented here. The Phoenix code, a sophisticated finite difference procedure, can handle far more complex situations than the elementary algorithm herein. These include evaporators with very small ratios of length to diameter and heat pipes of complex cross section.

The size of computer required for the Phoenix code and its cost make it less suitable for everyday heat pipe analysis. Furthermore, the simple
Figure 6. - Comparison of predicted values with Ivanovskii data.

Figure 7. - Comparison of predicted values with Ivanovskii data and results of other methods.

Equations derived from first principles in the present paper may facilitate a better physical understanding of the governing phenomena.

Concluding Remarks

The heat pipe vapor flow algorithm discussed herein has been shown to model the compressible vapor dynamics in the evaporator and adiabatic sections of pipes of conventional proportions. The model has introduced complexities which have usually been ignored. These include chemical equilibrium in the case of liquid metals, a velocity profile parameter dependent upon pressure and temperature, and a mean vapor temperature differing from the vapor interface temperature. Future work might be expended profitably on the introduction of a reasonable temperature profile into the energy equation, and the inclusion of droplet nucleation rates into the momentum and energy equations. Also, a satisfactory representation of the vapor effects in the condenser requires that the transition to turbulence, the treatment of condensation shocks, flow
recirculation, and other phenomena be modeled concisely for use in simple programs on small computers.

REFERENCES


## Abstract

This paper presents a heat pipe vapor flow algorithm suitable for use in codes on microcomputers. The incompressible heat pipe vapor flow studies of Busse are extended to incorporate compressibility effects. The Busse velocity profile factor is treated as a function of temperature and pressure. The assumption of a uniform saturated vapor temperature determined by the local pressure at each cross section of the pipe is not made. Instead, a mean vapor temperature, defined by an energy integral, is determined in the course of the solution in addition to the pressure, saturation temperature at the wall, and the Busse velocity profile factor. For alkali metal working fluids, local species equilibrium is assumed. Temperature and pressure profiles are presented for several cases involving sodium heat pipes. An example for a heat pipe with an adiabatic section and two evaporators in sequence illustrates the ability to handle axially varying heat input. A sonic limit plot for a short evaporator falls between curves for the Busse and Levy inviscid sonic limits.

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