

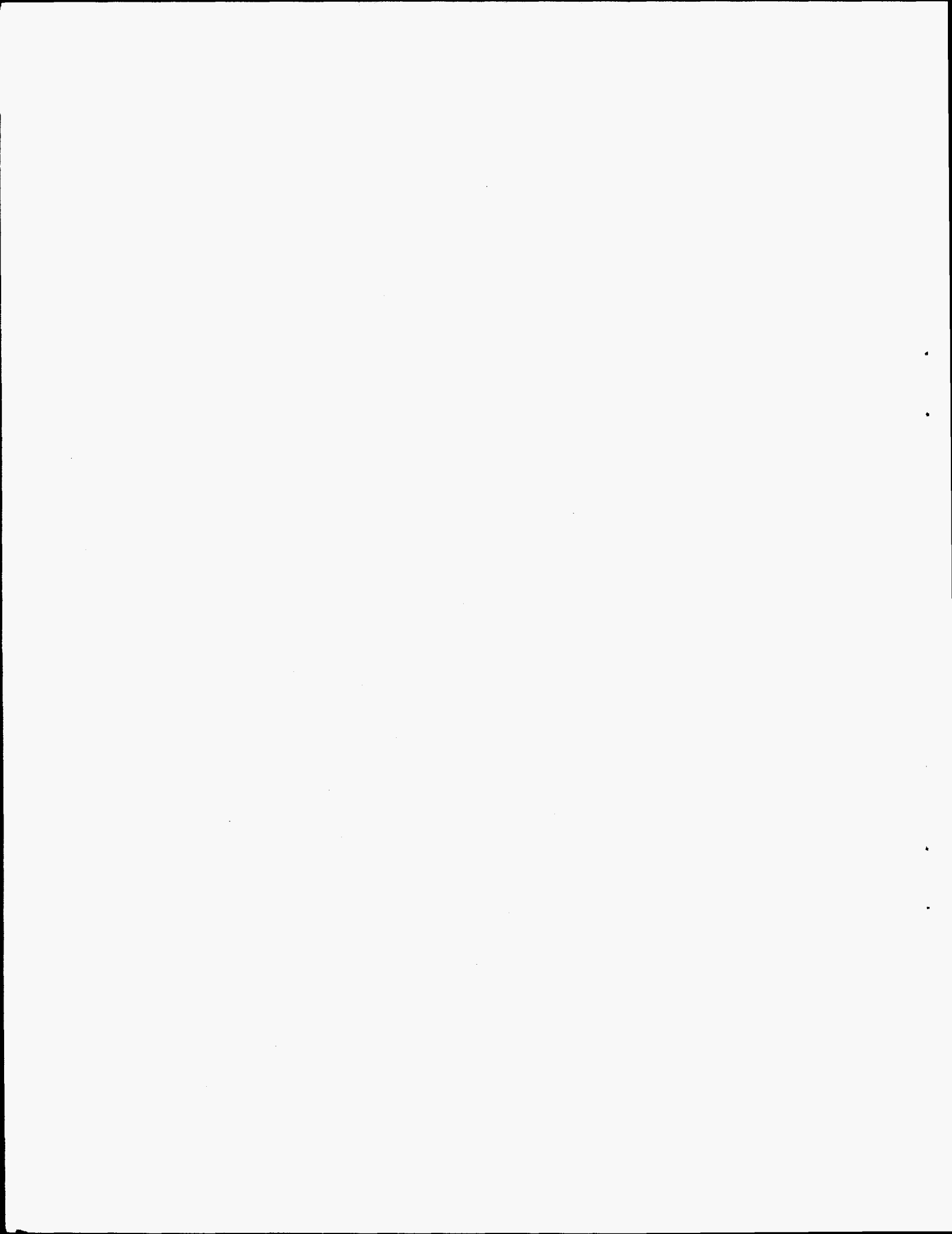
# **SCDAP/RELAP5 Modeling of Heat Transfer and Flow Losses in Lower Head Porous Debris**

**L. J. Siefken  
E. W. Coryell  
S. Paik**

**Published September 1998**

**Idaho National Engineering and Environmental Laboratory  
Lockheed Martin Idaho Technologies Company  
Idaho Falls, Idaho 83415**

**Prepared for the  
U.S. Department of Energy  
Assistant Secretary for  
Environmental Management  
Under DOE Idaho Operations Office  
Contract DE-AC07-94ID13223**



## **DISCLAIMER**

**This report was prepared as an account of work sponsored by an agency of the United States Government. Neither the United States Government nor any agency thereof, nor any of their employees, make any warranty, express or implied, or assumes any legal liability or responsibility for the accuracy, completeness, or usefulness of any information, apparatus, product, or process disclosed, or represents that its use would not infringe privately owned rights. Reference herein to any specific commercial product, process, or service by trade name, trademark, manufacturer, or otherwise does not necessarily constitute or imply its endorsement, recommendation, or favoring by the United States Government or any agency thereof. The views and opinions of authors expressed herein do not necessarily state or reflect those of the United States Government or any agency thereof.**

## **DISCLAIMER**

**Portions of this document may be illegible in electronic image products. Images are produced from the best available original document.**



## Abstract

Designs are described for implementing models for calculating the heat transfer and flow losses in porous debris in the lower head of a reactor vessel. The COUPLE model in SCDAP/RELAP5 represents both the porous and nonporous debris that results from core material slumping into the lower head. Currently, the COUPLE model has the capability to model convective and radiative heat transfer from the surfaces of nonporous debris in a detailed manner and to model only in a simplistic manner the heat transfer from porous debris. In order to advance beyond the simplistic modeling for porous debris, designs are developed for detailed calculations of heat transfer and flow losses in porous debris. Correlations are identified for convective heat transfer in porous debris for the following modes of heat transfer; (1) forced convection to liquid, (2) forced convection to gas, (3) nucleate boiling, (4) transition boiling, and (5) film boiling. Interphase heat transfer is modeled in an approximate manner. A design is also described for implementing a model of heat transfer by radiation from debris to the interstitial fluid. A design is described for implementation of models for flow losses and interphase drag in porous debris. Since the models for heat transfer and flow losses in porous debris in the lower head are designed for general application, a design is also described for implementation of these models to the analysis of porous debris in the core region. A test matrix is proposed for assessing the capability of the implemented models to calculate the heat transfer and flow losses in porous debris. The implementation of the models described in this report is expected to improve the COUPLE code calculation of the temperature distribution in porous debris and in the lower head that supports the debris. The implementation of these models is also expected to improve the calculation of the temperature and flow distribution in porous debris in the core region.



# CONTENTS

ABSTRACT .....	iii
FIGURES .....	vi
TABLES .....	vi
1. INTRODUCTION .....	1
2. REVIEW OF MODELS FOR HEAT TRANSFER BETWEEN POROUS DEBRIS AND INTERSTITIAL FLUID .....	2
2.1 Single Phase Vapor Regime .....	3
2.2 Single Phase Liquid Regime .....	7
2.3 Debris Heat Transfer for Two-Phase Flow .....	9
2.3.1 Nucleate Boiling .....	9
2.3.2 Film Boiling .....	11
2.3.3 Transition Boiling .....	13
2.3.4 Transition from Film Boiling to Convection to Steam .....	13
2.3.5 Radiation Heat Transfer .....	14
2.3.6 Total Heat Transfer to Liquid and Vapor Phases for Two-Phase Flow .....	15
2.4 Interphase Heat Transfer .....	17
3. IMPLEMENTATION OF CONVECTIVE AND RADIATIVE HEAT TRANSFER MODELS INTO COUPLE .....	17
4. FLOW LOSSES AND INTERPHASE DRAG IN POROUS DEBRIS .....	26
4.1 Flow regime identification .....	26
4.2 Models for Flow Losses .....	31
4.2.1 Drag Force for Superheated Steam (Single-Phase) .....	31
4.2.2 Drag Force for Subcooled and Saturated Liquid (Single-Phase) .....	31
4.2.3 Drag Force for Two-Phase Flow .....	32
4.2.3.1 Particle-Gas Drag Force for Two-Phase Flow .....	32
4.2.3.2 Particle-Liquid Drag Force for Two-Phase Flow .....	33
4.3 Models for interphase drag .....	34
5. IMPLEMENTATION OF MODELS FOR FLOW LOSSES AND INTERPHASE DRAG .....	36
5.1 Single Phase .....	37
5.2 Two Phase .....	38
6. EXTENSIONS TO IN-CORE POROUS DEBRIS .....	42
7. ASSESSMENT OF IMPLEMENTED MODELS .....	42
8. SUMMARY .....	43
9. REFERENCES .....	44

## FIGURES

1-1.	Schematic of situation represented by the COUPLE model (particle size of debris greatly exaggerated).....	1
4-1.	Schematic of pre-surface dryout flow regimes. ....	27
4-2.	Schematic of post-surface dryout flow regimes. ....	28

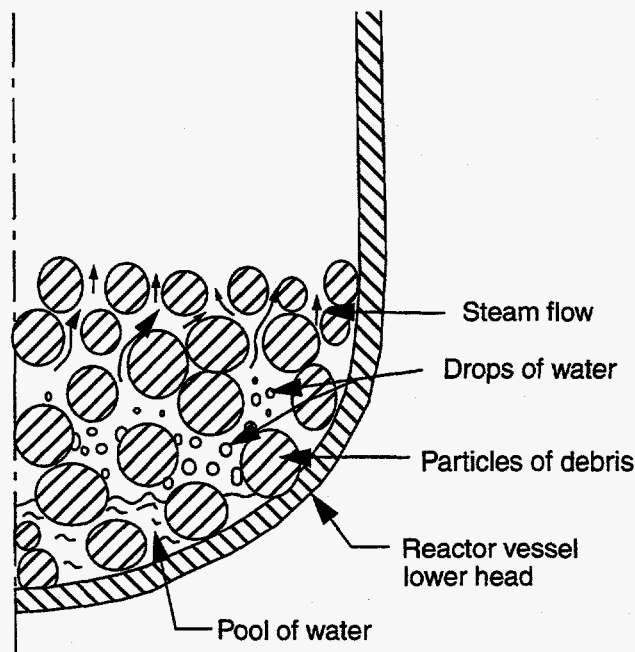
## TABLES

2-1.	Regimes of convective heat transfer and corresponding range in values of volume fraction of liquid and debris temperature. ....	2
2-2.	Definition of symbols in Table (2-1).....	3
3-1.	Variables added to COUPLE data base for modeling heat transfer to fluid in open porosity. ....	18
3-2.	Modifications of subroutine EGEN2 for modeling convective and radiative cooling.....	19
3-3.	Variables in common block debcom that are input and output variables for subroutine HTRC3B for porous debris heat transfer. ....	23
3-4.	Fortran modifications to subroutine HTRC3B for porous debris heat transfer.....	23
3-5.	Basic structure of subroutine HTRC3B for calculating debris to fluid heat transfer.....	24
3-6.	Fortran changes in subroutine COUPLE for implementing new models for heat transfer in porous debris.....	25
3-7.	Fortran lines added to RELAP5 subroutine PHANTV to model interphase heat transfer in porous debris.....	26
5-1.	Modification of RELAP5 subroutine HLOSS for modeling of flow losses in porous debris.....	39
6-1.	Modifications of SCDAP subroutine SCDAD5 for application to in-core debris of detailed models for cooling of porous debris.....	42
7-1.	Matrix of test problems for assessing models for cooling of porous debris.....	43

# 1. Introduction

A bed of porous debris may accrete in the lower head of a reactor vessel during a severe accident. If the porous debris is deep and covered with water, then a calculation needs to be made to determine whether the debris bed will locally dry out and heat up in spite of being covered with water. If the debris bed is hot and dry, then the reflood of the debris bed results in a quenching process that needs to be modeled to calculate the transient cooling of the debris bed.

The COUPLE model in SCDAP/RELAP5<sup>1</sup> is intended calculate the heatup of the lower head and the debris that it supports. The situation that the COUPLE model is designed to represent is shown in Figure 1-1. Currently, the COUPLE model has the capability to model convective and radiative heat transfer from the surfaces of nonporous debris in a detailed manner and to model only in a simplistic manner the heat transfer from porous debris. If water is present anywhere in porous debris, the entire porous debris bed is assumed to be quenched and all of the debris bed heat generation is transferred to boiling of water, which results in a reduction of the amount of water inside the porous debris. After all of the water has boiled off, then the debris bed is assumed to be cooled only at its outer boundaries. In order to advance beyond these simplifications in modeling, designs are developed for a detailed calculation of the heat transfer and flow losses in porous debris. This report describes these designs and their implementation into SCDAP/RELAP5.



**Figure 1-1.** Schematic of situation represented by the COUPLE model (particle size of debris greatly exaggerated).

This report is organized as follows. Section 2 identifies the correlations and models for convective and radiative heat transfer to be implemented into the COUPLE model. These correlations have been

previously compiled<sup>2</sup> and subjected to peer review.<sup>3</sup> Section 3 describes the basic features for implementation of the models for convective and radiative heat transfer. Models for flow losses and interphase heat transfer are described in Section 4 and the implementation of these models is described in Section 5. The implementation of the models developed for analysis of porous debris in the lower head results in a straightforward application of these models to the analysis of porous debris in the core region. This application is described in Section 6. Section 7 presents plans for testing and assessing the implemented models. A summary of the extensions in modeling proposed for SCDAP/RELAP5 is given in Section 8. The references are listed in Section 9.

## 2. Review of Models for Heat Transfer between Porous Debris and Interstitial Fluid

This section reviews models for convective and radiative heat transfer that have been previously published and subjected to peer review,<sup>2,3</sup> and which are recommended for incorporation into the COUPLE model. Seven regimes of convective heat transfer are identified and correlations for heat transfer are obtained for each regime. Equations are also presented for calculating the radiative heat transfer between porous debris and the interstitial fluid.

The regimes of convective heat transfer are distinguished by the values of two parameters; (1) volume fraction of liquid in the open porosity of the coolant, and (2) temperature of the debris. The regimes of heat transfer range from nucleate boiling in two-phase coolant to natural convection in steam. The various regimes of convective heat transfer and the corresponding ranges in values of volume fraction of liquid and debris temperature are identified in Table 2-1. The symbols used in Table 2-1 are defined in Table 2-2. In Table 2-1 several temperature thresholds are identified for transition from one regime of heat transfer to another regime of heat transfer. One of these temperature thresholds, namely  $T_{sat}$ , is determined by the water properties package for SCDAP/RELAP5. Another threshold temperature, namely  $T_{TF}$ , is determined from experimental results. The other temperature thresholds are determined by matching a heat flux from one regime of heat transfer with the heat flux from the interfacing regime of heat transfer.

**Table 2-1.** Regimes of convective heat transfer and corresponding ranges in values of volume fraction of liquid and debris temperature.

Phase state of fluid	Mode of heat transfer	Range of void fraction of vapor	Range of debris temperature (K)
<i>single phase vapor</i>	<i>forced convection and natural convection</i>	1.0	$T_D > T_{sat}$
<i>single phase liquid</i>	<i>forced convection and natural convection</i>	0.0	$T_D \leq T_{sat}$
<i>two-phase</i>	<i>nucleate boiling</i>	$0.0 \leq \alpha_g < 1.0$	$T_{CN} < T_D < T_{nuc}$
<i>two-phase</i>	<i>transition boiling</i>	$0.0 \leq \alpha_g < 1.0$	$T_{nuc} < T_D < T_{TF}$
<i>two-phase</i>	<i>film boiling</i>	$\alpha_g \leq \alpha_4$	$T_D > T_{TF}$
<i>two-phase</i>	<i>transition from film boiling to convection to vapor</i>	$\alpha_4 \leq \alpha_g < 1.0$	$T_D > T_{TF}$

**Table 2-2.** Definition of symbols in Table 2-1.

Symbol	Units	Definition
$\alpha_g$	--	<i>volume fraction of vapor in fluid</i>
$T_D$	K	<i>temperature of debris</i>
$T_{sat}$	K	<i>saturation temperature of fluid</i>
$T_{CN}$	K	<i>temperature of debris at which heat flux using convection correlation equals heat flux using nucleate boiling correlation</i>
$T_{nuc}$	K	<i>temperature of debris at which heat flux using nucleate boiling correlation equals critical heat flux</i>
$T_{TF}$	K	<i>temperature of debris at which transition boiling heat transfer ends and film boiling heat transfer begins</i>
$\alpha_4$	--	<i>void fraction at which flow regime changes from inverted slug-mist flow to mist flow (~ 0.925).</i>

## 2.1 Single Phase Vapor Regime

The heat transfer correlation developed by Tung<sup>4</sup> will be used to calculate the debris-to-vapor convective heat transfer. In this correlation, the Nusselt number is given by the equation

$$Nu_{conv} = 0.27Re^{0.8}Pr^{0.4} \quad (2-1)$$

where

$Nu_{conv}$  = Nusselt number for convection,

$Re$  = Reynold's number,

$Pr$  = Prandtl number.

The Reynold's number is given by the equation

$$Re = \rho_g V_g D_p / \mu_g \quad (2-2)$$

where

$\rho_g$  = density of vapor ( $kg/m^3$ ),

- $V_g$  = velocity of vapor (m/s),  
 $D_p$  = effective diameter of debris particle (m),  
 $\mu_g$  = viscosity of vapor (kg/m · s).

The Prandtl number is given by the equation

$$Pr = \mu_g c_g / k_g \quad (2-3)$$

where

- $\mu_g$  = viscosity of vapor (kg/m · s),  
 $c_g$  = heat capacity of vapor (J/kg · K),  
 $k_g$  = thermal conductivity of vapor (W/m · K).

The ranges of parameters for which this correlation is based are

$$0.7 \leq Pr \leq 5$$

$$18 \leq Re \leq 2400$$

$$\varepsilon \approx 0.4.$$

For the case of low fluid velocity, the Nusselt number for natural convection will also be calculated. If the Nusselt number for natural convection is greater than that for forced convection, then the natural convection Nusselt number will be applied. The natural convection Nusselt number is taken from Tung<sup>4</sup> as

$$Nu_{nat} = KRa^{0.25} \quad (2-4)$$

where

$Nu_{nat}$  = Nusselt number for natural convection,

$$K = \begin{cases} 0.3 & 0 \leq Ra \leq 50 \\ 0.4 & 50 \leq Ra \leq 200 \\ 0.5 & 200 \leq Ra \leq 10^6 \\ 0.6 & 10^6 \leq Ra \leq 10^8 \end{cases}$$

$Ra$  = Rayleigh number.



The Rayleigh number is calculated by the equation

$$Ra = Gr \cdot Pr = \frac{\rho_g^2 g D_p^3 \beta \Delta T}{\mu_g^2} Pr \quad (2-5)$$

where

- g = acceleration of gravity (m/s<sup>2</sup>),  
 β = volume coefficient of expansion of vapor (1/K),  
 ΔT = temperature difference between debris and vapor (T<sub>D</sub> - T<sub>g</sub>).

The heat transferred to the vapor by convection will be calculated by the equation

$$Q_{conv} = A_s \max(Nu_{conv}, Nu_{nat}) \frac{k_g}{D_p} (T_D - T_g) \quad (2-6)$$

where

- Q<sub>conv</sub> = heat transferred to vapor by convection (W/m<sup>3</sup>),  
 A<sub>s</sub> = surface area of debris per unit volume (m<sup>2</sup>/m<sup>3</sup>),  
 T<sub>D</sub> = temperature of debris particles (K),  
 T<sub>g</sub> = temperature of vapor (K).

The surface area of debris per unit volume is calculated by the equation

$$A_s = \frac{6(1 - \epsilon)}{D_p} \quad (2-7)$$

where

- ε = porosity of debris.

The heat transferred to the vapor by radiation is calculated by the equation

$$Q_{rad} = A_s F_g \sigma (T_D^4 - T_g^4), \quad (2-8)$$

where

- $Q_{\text{rad}}$  = heat transferred to vapor by radiation (W/m<sup>3</sup>),  
 $F_g$  = gray-body factor,  
 $\sigma$  = Stefan-Boltzmann constant ( $5.668 \times 10^{-8}$  W/m<sup>2</sup>K<sup>4</sup>).

The gray body factor is calculated by the equation

$$F_g = 1/[R_1(1 + R_3/R_1 + R_3/R_2)] \quad (2-9)$$

where

- $R_1$  =  $(1 - \epsilon_g)/\epsilon_g$ ,  
 $R_2$  =  $1/\epsilon_g$ ,  
 $R_3$  =  $1 + (1 - \epsilon_w)/\epsilon_w$ ,  
 $\epsilon_g$  =  $1 - \exp(-a_g L_m)$ ,  
 $\epsilon_w$  = 0.7,  
 $a_g$  = absorption coefficient for vapor,  
 $L_m$  = mean path length (m).

The absorption coefficient,  $a_g$  is calculated by the SCDAP subroutine EMISSV.<sup>1</sup> An estimation of the mean path is obtained by assuming it equal to the hydraulic diameter.<sup>5</sup> The hydraulic diameter is estimated by the equation

$$L_m = \frac{4\epsilon D_p}{6(1 - \epsilon)} \quad (2-10)$$

The total heat transfer to the vapor is then

$$Q = Q_{\text{conv}} + Q_{\text{rad}} \quad (2-11)$$

where

- $Q$  = total heat transfer to vapor (W/m<sup>3</sup>).

## 2.2 Single Phase Liquid Regime

The heat transfer correlation presented by Gunn<sup>6</sup> is used to calculate the volumetric heat transfer coefficient for the covered regime. This correlation is applicable for water that is either subcooled or saturated. The correlation for the Nusselt number is given by the equation

$$Nu = (7 - 10\varepsilon + 5\varepsilon^2)(1 + 0.7Re^{0.3}Pr^{0.333}) + (1.33 - 2.4\varepsilon + 1.2\varepsilon^2)Re^{0.7}Pr^{0.333} \quad (2-12)$$

where

Pr = Prandtl number.

The Prandtl number is calculated by the equation

$$Pr = \mu_f c_f / k_f \quad (2-13)$$

where

$\mu_f$  = viscosity of the liquid (kg/m · s),

$c_f$  = heat capacity of the liquid (J/kg · K),

$k_f$  = thermal conductivity of the water (W/m · K).

The Reynold's number is calculated by the equation

$$Re = v_f \rho_f l / \mu_f \quad (2-14)$$

where

$v_f$  = velocity of the liquid (m/s),

$\rho_f$  = density of the liquid (kg/m<sup>3</sup>),

$l$  = characteristic length as defined below (m).

The volumetric heat transfer coefficient is calculated by the equation

$$h_v = Nu \frac{k_f}{l^2} \quad (2-15)$$

The characteristic length is calculated by the equation

$$l = b/a \quad (2-16)$$

where

- b = inertial coefficient in Kozeny-Carman equation (1/m),  
a = viscous coefficient in Kozeny-Carman equation (1/m<sup>2</sup>).

The coefficients b and a are calculated by the equations

$$b = 1.75(1 - \epsilon)/\epsilon^3 D_p, \quad (2-17)$$

$$a = 150(1 - \epsilon)^2/\epsilon^3 D_p^2$$

where

- $\epsilon$  = porosity of debris,  
 $D_p$  = effective diameter of debris particles (m).

The total heat transfer to the fluid is calculated by the equation

$$Q = h_v(T_D - T_f) \quad (2-18)$$

where

- Q = total heat transfer to the fluid (W/m<sup>3</sup>),  
 $T_D$  = surface temperature of debris (K),  
 $T_f$  = temperature of liquid (K).

The heat transfer to the vapor phase and the volumetric vapor generation rate are equal to zero for this heat transfer regime.

The forced convection and natural convection heat transfer to the liquid phase can also be calculated by Equations (2-1) and (2-6) with the properties of the liquid phase substituted in place of properties of the vapor phase.<sup>3</sup> It is recommended that a user-defined selection between the two correlations be maintained initially and an assessment performed to evaluate which of the two equations produces the best results.<sup>3</sup>

## 2.3 Debris Heat Transfer for Two-Phase Flow

The debris-to-fluid heat transfer in the two-phase region is a complex process that will be represented in the simplified manner devised by Tutu, et al,<sup>7</sup> except for modifications made to make the heat transfer modeling consistent with the flow regime modeling. In view of the fact that there is an absence of experimental data and theoretical models for local heat transfer coefficients for two-phase conditions, a simplified approach is required.

Four modes of convective heat transfer are considered: (1) nucleate boiling; (2) film boiling; (3) transition boiling; and (4) transition from film boiling to convection to vapor. The mode of heat transfer that is in effect is a function of the debris temperature and the volume fraction of vapor in the fluid. The range of conditions for each mode of heat transfer have been summarized in Table 2-1. The symbols in Table 2-1 have been defined in Table 2-2. Heat transfer by radiation is also taken into account.

### 2.3.1 Nucleate Boiling

The heat transfer coefficient for nucleate boiling will be calculated by a correlation for pool boiling that was developed by Rohsenow<sup>7</sup> and used by Tutu, et al.<sup>15</sup> This correlation is

$$h_{snuc} = 4.63 \times 10^6 f(\text{prop})(T_D - T_{sat})^m \quad (2-19)$$

where

$h_{snuc}$  = heat transfer coefficient for nucleate boiling mode of heat transfer ( $W/m^2 \cdot K$ ),

$f(\text{prop})$  = function that is combination of fluid properties as defined below,

$m$  = exponent that is function of particle diameter as shown below.

The function of fluid properties is

$$f(\text{prop}) = \frac{\mu_f c_{pf}^3}{(h_{fg}^2) \left[ \frac{\sigma}{g(\rho_f - \rho_g)} \right]^{0.5} \left( \frac{c_{pf} \mu_f}{k_f} \right)^{4.913}} \quad (2-20)$$

where

$\mu_f$  = viscosity of liquid water ( $kg/m \cdot s$ ),

$c_{pf}$  = heat capacity of liquid water ( $J/kg \cdot K$ ),

$h_{fg}$  = latent heat of vaporization ( $J/kg \cdot K$ ),

- $\sigma$  = surface tension ( $\text{kg/s}^2$ ),  
 $g$  = acceleration of gravity ( $9.8 \text{ m/s}^2$ ),  
 $\rho_f$  = density of saturated liquid ( $\text{kg/m}^3$ ),  
 $\rho_g$  = density of saturated liquid ( $\text{kg/m}^3$ ),  
 $k_f$  = thermal conductivity of liquid water ( $\text{W/m}\cdot\text{K}$ ).

The exponent  $m$  is calculated by the equation<sup>3</sup>

$$m = 3.3 + 9.0e^{-d} \quad (2-21)$$

where

$$d = \left\{ \frac{D_p}{\left[ \frac{\sigma}{g(\rho_f - \rho_g)} \right]^{0.5}} \right\}^{0.5}$$

If  $d$  is calculated to be less than 1.4142, then  $d$  is set to a value of 1.4142.

The debris temperature at which the transition from the forced convection to single phase liquid mode of heat transfer to the nucleate boiling mode of heat transfer takes place is equal to  $T_{\text{CN}}$ , which is determined by solving the following equation for  $T_{\text{CN}}$ :

$$h_v(T_{\text{CN}} - T_{\text{sat}}) = 4.63 \times 10^6 f(\text{prop})(T_{\text{CN}} - T_{\text{sat}})^m A_s \quad (2-22)$$

where

- $h_v$  = volumetric heat transfer coefficient for forced convection to liquid,  
 $T_{\text{CN}}$  = temperature of debris at which heat flux using forced convection correlation equals heat flux using nucleate boiling correlation (K),  
 $A_s$  = surface area of particles as defined by Equation (2-7) ( $\text{m}^2/\text{m}^3$ ).

The maximum temperature of debris for the nucleate boiling mode of heat transfer is determined by solving the following equation for  $T_{\text{nuc}}$ :

$$4.63 \times 10^6 f(\text{prop})(T_{\text{nuc}} - T_{\text{sat}})^m (T_{\text{nuc}} - T_{\text{sat}}) = q_{\text{CHF}} \quad (2-23)$$

where

- $T_{\text{nuc}}$  = maximum particle temperature for nucleate boiling mode of heat transfer (K),  
 $q_{\text{CHF}}$  = critical heat flux calculated as shown below ( $\text{W}/\text{m}^2$ ).

The critical heat flux is calculated by a correlation for spheres developed by Ded and Lienhard:<sup>8</sup>

$$q_{\text{CHF}} = 0.11 F_d h_{fg} \rho_g [g \sigma ((\rho_f - \rho_g) / \rho_g^2)]^{0.25} \quad (2-24)$$

where

- $F_d$  = factor correcting for debris particle size as defined below,  
 $h_{fg}$  = latent heat of vaporization (J/kg),  
 $g$  = acceleration of gravity ( $9.8 \text{ m}/\text{s}^2$ ),  
 $\rho_f$  = density of saturated liquid ( $\text{kg}/\text{m}^3$ ),  
 $\rho_g$  = density of saturated liquid ( $\text{kg}/\text{m}^3$ ),  
 $\sigma$  = surface tension ( $\text{kg}/\text{s}^2$ ).

The factor  $F_d$  is calculated by the equation<sup>3</sup>

$$F_d = 1 - 3.8e^{-d} \quad (2-25)$$

where

$$d = \left\{ \frac{D_p}{\left[ \frac{\sigma}{g(\rho_f - \rho_g)} \right]^{0.5}} \right\}^{0.5}$$

If  $d$  is calculated to be less than 1.4142, then  $d$  is set to a value of 1.4142.

### 2.3.2 Film Boiling

The correlation developed by Dhir and Lienhard<sup>9</sup> will be used to calculate the surface heat transfer coefficient for the film boiling mode of heat transfer. According to their correlation, the Nusselt number is calculated by the equation

$$Nu_{fb} = 0.67 \left[ \frac{\rho_g (\rho_f - \rho_g) g h_{fgs} D_p^3}{\mu_g k_g \Delta T} \right]^{0.25} \quad (2-26)$$

where

$Nu_{fb}$	=	Nusselt number for film boiling mode of heat transfer,
$\rho_g$	=	density of saturated vapor (kg/m <sup>3</sup> ),
$\rho_f$	=	density of saturated liquid (kg/m <sup>3</sup> ),
$g$	=	acceleration of gravity (9.8 m/s <sup>2</sup> ),
$D_p$	=	effective diameter of particles (m),
$\mu_g$	=	viscosity of saturated vapor (kg/m s),
$k_g$	=	thermal conductivity of saturated vapor (W/m K),
$\Delta T$	=	$T_g - T_{sat}$ ,
$T_g$	=	temperature of the vapor (K),
$T_{sat}$	=	saturation temperature (K),
$h_{fgs}$	=	$h_{fg} + 0.68 c_{pg} \Delta T$ ,
$c_{pg}$	=	heat capacity of water vapor (J/kg K),
$h_{fg}$	=	latent heat of vaporization (J/kg).

This correlation was developed using polished spheres. Experimental results indicate that for particles with an oxide layer on the surface and a low superheat, the Nusselt number may be 80% higher than that for particles with a polished surface.<sup>4</sup> At high superheats, the heat transfer coefficients for oxidized and polished particles converge. Although particles in a debris bed in a nuclear reactor are expected to be oxidized, they may also be very hot, so a multiplier to account for oxidized surfaces will not be applied.

The surface heat transfer coefficient for film boiling is calculated by the equation

$$h_{sfb} = (Nu_{fb}) \frac{k_g}{D_p} \quad (2-27)$$



where

$h_{sfb}$  = surface heat transfer coefficient for film boiling ( $W/m^2 \cdot K$ ).

### 2.3.3 Transition Boiling

In the transition boiling mode of heat transfer, when the debris temperature is between  $T_{nuc}$  and  $T_{TF}$ , the heat transfer coefficient is calculated by the equation

$$h_{str} = \frac{(T_D - T_{nuc})}{(T_{TF} - T_{nuc})} [h_{sfb} - h_{snuc}] + h_{snuc} \quad (2-28)$$

where

$h_{str}$  = heat transfer coefficient for transition boiling mode of heat transfer ( $W/m^2 \cdot K$ ),

$T_{TF}$  = temperature of debris at which transition boiling heat transfer ends and film boiling heat transfer begins.

The variable  $T_{TF}$  is calculated by the equation<sup>3</sup>;

$$T_{TF} = 0.16 \frac{\rho_g h_{fg}}{k_g} \left[ \frac{g(\rho_f - \rho_g)}{(\rho_f + \rho_g)} \right]^{2/3} \left[ \frac{\mu_g}{g(\rho_f + \rho_g)} \right]^{1/3} \left[ \frac{\sigma}{g(\rho_f + \rho_g)} \right]^{1/2} + T_{sat} \quad (2-29)$$

### 2.3.4 Transition from Film Boiling to Convection to Steam

The transition from the film boiling mode of heat transfer to the convection to steam mode of heat transfer is assumed to occur when the void fraction of vapor is between  $\alpha_4$  and 1. The transition from the inverted slug-mist flow regime to the mist flow regime occurs at a void fraction  $\alpha_4$ . The equation for calculating  $\alpha_4$  is described in Reference 2. In this range of void fractions, the heat transfer to the liquid and vapor phases are calculated by the equations

$$Q_{cf} = (1 - W_{fg}) A_s h_{sfb} (T_D - T_{sat}) \quad (2-30)$$

$$Q_{cg} = W_{fg} Q_{conv}$$

where

$Q_{cf}$  = total heat transfer to liquid phase by convection ( $W/m^3$ ),

$Q_{cg}$  = total heat transfer to vapor phase by convection ( $W/m^3$ ),

$W_{fg}$  = weighting function as defined below,

$Q_{conv}$  = heat transfer to vapor as calculated by Equation (2-6) of Section 2.1 ( $W/m^3$ ).

The weighting function will be<sup>4</sup>

$$W_{fg} = y^2(3 - 2y) \quad (2-31)$$

where

$W_{fg}$  = weighting function for interpolation between film boiling and convection to vapor modes of heat transfer,

$$y = \frac{\alpha_g - \alpha_4}{1 - \alpha_4},$$

$\alpha_g$  = volume fraction of vapor,

$\alpha_4$  = void fraction at which flow regime changes from inverted slug-mist flow to mist flow.

### 2.3.5 Radiation Heat Transfer

The heat transferred to the liquid and vapor phases by radiation is calculated by the equations

$$Q_{rf} = A_s F_f \sigma (T_D^4 - T_{sat}^4) \quad (2-32)$$

$$Q_{rg} = A_s F_g \sigma (T_D^4 - T_g^4)$$

where

$Q_{rf}$  = heat transferred to the liquid phase by radiation ( $W/m^3$ ),

$Q_{rg}$  = heat transferred to the vapor phase by radiation ( $W/m^3$ ),

$F_f$  = gray body factor for liquid,

$F_g$  = gray body factor for vapor,

$\sigma$  = Stefan-Boltzmann constant ( $5.668 \times 10^{-8} W/m^2K^4$ ),

$T_g$  = temperature of vapor phase (K).

The gray body factors are defined as

$$F_f = 1/[R_2(1 + R_3/R_1 + R_3/R_2)], \quad (2-33)$$

$$F_g = 1/[R_1(1 + R_3/R_1 + R_3/R_2)].$$

The R terms are given as

$$R_1 = (1 - \epsilon_g)/[\epsilon_g(1 - \epsilon_g/\epsilon_f)], \quad (2-34)$$

$$R_2 = (1 - \epsilon_f)/[\epsilon_g(1 - \epsilon_g/\epsilon_f)],$$

$$R_3 = 1/(1 - \epsilon_g\epsilon_f) + (1 - \epsilon_w)/\epsilon_w.$$

The emissivities,  $\epsilon$ , are given as

$$\epsilon_g = 1 - \exp(-a_g L_m), \quad (2-35)$$

$$\epsilon_f = 1 - \exp(-a_f L_m),$$

$$\epsilon_w = 0.7$$

$L_m$  = mean path length.

The absorption coefficients,  $a_g$  and  $a_f$ , are calculated by subroutine EMISSV.<sup>1</sup> The path length  $L_m$  is calculated by Equation (2-10).

### 2.3.6 Total Heat Transfer to Liquid and Vapor Phases for Two-Phase Flow

The heat transfer to the liquid phase by convection is calculated by the equation

$$Q_{cf} = A_s h_{sf}(T_D - T_{sat}) \quad (2-36)$$

where

$Q_{cf}$  = total heat transfer to liquid phase by convection ( $W/m^3$ ),

$A_s$  = surface area of particles as calculated by Equation (2-7) for modes other than forced convection to liquid ( $m^2/m^3$ ). For forced convection to liquid mode,  $A_s$

equals 1 (unitless).

$h_{sf}$  = heat transfer coefficient corresponding with the applicable mode of heat transfer, as defined in Table (2-1) for modes other than forced convection to liquid ( $W/m^2 \cdot K$ ). For forced convection to liquid, units of  $h_{sf}$  are ( $W/m^3 \cdot K$ ).

The total heat transferred to the fluid is calculated by the equation

$$Q_{tot} = Q_{cf} + Q_{rf} + Q_{cg} + Q_{rg} \quad (2-37)$$

where

$Q_{tot}$  = total heat transferred to the fluid (vapor and liquid phase) ( $W/m^3$ ). This variable, when multiplied by the volume of a control volume corresponds with the variable  $Q$  in the RELAP5 code.

If the void fraction of vapor is less than  $\alpha_4$ , the terms for heat transfer to the vapor in the above equation are equal to zero.

The total heat transferred to the vapor phase is calculated by the equation

$$Q_{totg} = Q_{cg} + Q_{rg} \quad (2-38)$$

where

$Q_{totg}$  = total heat transferred to the vapor phase ( $W/m^3$ ). This variable, when multiplied by the volume of a control volume, corresponds with the variable  $Q_{WG}$  in the RELAP5 code. If the void fraction of vapor is less than  $\alpha_4$ , this term is equal to zero.

The vapor generation is calculated by the equation

$$\Gamma_w = (Q_{cf} + Q_{rf})/h_{fg} \quad (2-39)$$

where

$\Gamma_w$  = volumetric vapor generation rate ( $kg/m^3s$ ). This variable, when multiplied by the volume of porous debris in a control volume and then divided by the fluid volume of the control volume, corresponds with the variable  $GAMMAW$  in the RELAP5 code.

## 2.4 Interphase Heat Transfer

The heat transfer between the liquid and vapor phases of the fluid in porous debris will be modeled in an approximate manner. The thermal equilibrium model will be used. This model will be applied by setting the interfacial heat transfer coefficient to a large value for RELAP5 control volumes that contain porous debris. The suitability of this assumption will be assessed by comparing calculated and measured temperatures at the top of an initially hot debris bed that was quenched from the bottom. The temperature history in this region of the debris bed is strongly influenced by the temperature of the steam flowing through the debris bed. If the calculated and measured temperatures in this region of the debris bed are in good agreement, then the simplifying assumption of thermal equilibrium of the liquid and vapor phases is suitable.

## 3. Implementation of Convective and Radiative Heat Transfer Models into COUPLE

The cooling of porous debris can be treated as a heat transfer process that is parallel with the internal heat generation in the porous debris. This treatment is possible because the particles in the porous debris are generally very small compared with the size of a COUPLE node and thus the cooling is spatially and uniformly distributed through the node. The effect on debris heatup of convective and radiative cooling can be calculated by subtracting the amount of cooling per unit volume of debris from the amount of heat generation per unit volume of debris. The COUPLE code applies the heat generation term in calculating heat transport as follows;

$$(\rho C_v)_e \frac{\partial T}{\partial t} = \frac{\partial}{\partial x} \left( K_e \frac{\partial T}{\partial x} \right) + \frac{\partial}{\partial y} \left( K_e \frac{\partial T}{\partial y} \right) + Q \quad (3-1)$$

where

$$(\rho C_v)_e = P(\rho C_v)_l + (1 - P)(\rho C_v)_s$$

- $\rho$  = density ( $\text{kg/m}^3$ ),
- $C_v$  = constant volume specific heat,
- $K$  = thermal conductivity,
- $Q$  = volumetric heat generation rate ( $\text{W/m}^3$ ),
- $T$  = temperature,
- $P$  = porosity (pore volume/total volume),
- $e$  = equivalent,

l = liquid,  
s = solid.

To account for convective heat transfer in the debris, the heat generation term  $Q$  in Equation (3-1) will be replaced by the following term;

$$Q_{net} = Q - Q_c \quad (3-2)$$

where

$Q_c$  = heat transferred from debris to fluid by convective and radiative heat transfer ( $W/m^3$ ).

The variable  $Q_c$ , which is the volumetric rate of removal of heat by convective and radiative heat transfer, is calculated for every possible regime of heat transfer by the equations presented in Section 2.

The implementation of the models for convective and radiative heat transfer from the debris is accomplished by transforming the COUPLE Fortran variable that stores the volumetric heat generation into a variable that equals the term  $Q_{net}$  in Equation (3-2). The Fortran variable to be transformed is named  $bg(i)$ , which is the volumetric heat generation ( $W/m^3$ ) at node  $i$  of the COUPLE model mesh. This variable is calculated in subroutine EGEN2. After calculating  $bg(i)$  accounting for heat generation, a call will be made to a subroutine named HTRC3B to calculate the convective and radiative cooling at each node in the COUPLE mesh with debris. The COUPLE model data base will also be expanded to store the vapor generation rate and the heat transfer to the liquid and vapor phases of the coolant for each COUPLE node. The new variables to be added to the data base are defined in Table 3-1 and the fortran changes to be made to subroutine EGEN2 are described in Table 3-2.

**Table 3-1.** Variables added to COUPLE data base for modeling heat transfer to fluid in open porosity.

Fortran variable	Units	Definition
<i>ihpore(n)</i>	-	<i>indicator of whether n-th convective node has participated in heat transfer from porous debris to fluid, 0=no, 1=yes; it is defined in subroutine egen2 and stored in common block alcm</i>
<i>qcrdeb(i)</i>	W	<i>heat removed by convective and radiative heat transfer from debris at node i to fluid in open porosity at node i, array stored in common block alcm</i>
<i>iptpor</i>	-	<i>pointer to location of array ihpore in common block alcm, stored in common block iparm</i>
<i>iqcrdb</i>	-	<i>pointer to location of array qcrdeb in common block alcm, stored in common block iparm</i>

**Table 3-1.** Variables added to COUPLE data base for modeling heat transfer to fluid in open porosity. (continued)

Fortran variable	Units	Definition
<i>ndbhtr</i>	-	<i>indicator of option for modeling of debris to fluid heat transfer, 0=detailed model, 1=simplified model, stored in common block iparm</i>

**Table 3-2.** Modifications of subroutine EGEN2 for modeling convective and radiative cooling.

Line of Fortran	Comments	Line status <sup>a</sup>
<i>subroutine egen2(ix,gen,bg,xm2,ng2,ngnm,bv,engnod,powrat,</i>	<i>add arguments for variables needed in debris to fluid heat transfer</i>	<i>M</i>
<i># tz, pore, dimpe, qcoupl, qwgcou, gamcou, vole,</i>	<i>temperature, RELAP5 index pointer, porosity, particle size, total heat to fluid, heat to vapor, vapor generation rate, corner volumes,</i>	<i>N</i>
<i>qcrdeb, ir5vec, ihsave, ihpore)</i>	<i>heat removed, pointer to RELAP5 control volume indices, convection nodes, indicator of presence of porous debris</i>	<i>N</i>
<i>*in32 ir5vec,ihsave, ihpore</i>	<i>adjust for 32 bit integer in 64 bit word</i>	<i>M</i>
<i>*call comctl</i>	<i>add parameters needed for RELAP5 volume data block</i>	<i>N</i>
<i>*call voldat</i>	<i>RELAP5 volume data block</i>	<i>N</i>
<i>*call fast</i>	<i>RELAP5 data</i>	<i>N</i>
<i>*call scddat</i>	<i>add common block defining dimension of arrays in common block debcom</i>	<i>N</i>
<i>call debcom</i>	<i>add debcom common, which contains input and output variables for subroutine htrc3b</i>	<i>N</i>
<i>do 100 n=1,numel</i>	<i>numel=number of elements</i>	<i>E</i>

**Table 3-2.** Modifications of subroutine EGEN2 for modeling convective and radiative cooling. (continued)

Line of Fortran	Comments	Line status <sup>a</sup>
<i>i=ix(1,n)</i>	<i>identify nodes at each corner of element</i>	<i>E</i>
<i>nn=ndbreg+1</i>	<i>initial value for index nn, which is the second index in the two-dimensional arrays in common block debcom</i>	<i>N</i>
<i>mm=1</i>	<i>initial value for index mm, which is the first index in the two-dimensional arrays in common block debcom</i>	<i>N</i>
<i>.....</i>	<i>skip over coding</i>	<i>E</i>
<i>bg(i)=bg(i)+pwkinc*gen(1,n)/bv(i) + ...</i>	<i>bv(n) = volume of element, gen(1,n) = heat generation for first corner</i>	<i>E</i>
<i>if(pore(n).lt.0.1)go to 521</i>	<i>skip over node with nonporous debris</i>	<i>N</i>
<i>Note: Investigate possibility of calling htrc3b with mm=1 and nn=ndbreg for all COUPLE model nodes</i>		
<i>if(mm.lt.ndax)then</i>	<i>determine unique set of indices (mm,nn) for couple node i</i>	<i>N</i>
<i>mm=mm+1</i>		<i>N</i>
<i>else</i>		<i>N</i>
<i>nn=nn+1</i>		<i>N</i>
<i>if(nn.gt.ndxbrg)then</i>	<i>protect against possibility of overflowing dimension of debcom arrays</i>	<i>N</i>
<i>write message</i>		<i>N</i>
<i>stop</i>	<i>stop execution of code</i>	<i>N</i>
<i>end if</i>	<i>end of "if()then" block on protection</i>	<i>N</i>
<i>end if</i>	<i>end of "if()then" block on defining mm and nn indices</i>	<i>N</i>
<i>nvoldb(nn)=1</i>	<i>define one RELAP5 control volume for nn-th debris region</i>	<i>N</i>



**Table 3-2.** Modifications of subroutine EGEN2 for modeling convective and radiative cooling. (continued)

Line of Fortran	Comments	Line status <sup>a</sup>
<i>ir5=0</i>	<i>initialization</i>	<i>N</i>
<i>istop=0</i>	<i>initialization</i>	<i>N</i>
<i>do20ncon=1,ncev</i>	<i>begin do loop to identify RELAP5 volume that node i is connected to</i>	<i>N</i>
<i>ihpore(ncon)=0</i>	<i>set default value</i>	<i>N</i>
<i>if(istop.eq.1)go to 20</i>	<i>match already found</i>	<i>N</i>
<i>if(ihsave(ncon).eq.i.and.ntf2(ncon).ge.0)then</i>	<i>match found</i>	<i>N</i>
<i>ir5=ncon</i>	<i>store sequence number of convective node</i>	<i>N</i>
<i>ihpore(ncon)=1</i>	<i>identify that convective node is involved with heat transfer to interstitial fluid</i>	<i>N</i>
<i>istop=1</i>	<i>indicate that no further searching required to identify sequence number of convective node</i>	<i>N</i>
<i>end if</i>		<i>N</i>
<i>20 continue</i>		<i>N</i>
<i>if(ir5.eq.0)go to 521</i>	<i>ir5 = 0 = no porous debris at node</i>	<i>N</i>
<i>numdbv(mm,nn)=ir5vec(ncon)+filndx(4)</i>	<i>store RELAP5 volume index for debris location (mm,nn)</i>	<i>N</i>
<i>idbvol(ir5vec(ncon)+filndx(4))=2</i>	<i>identify that porous debris present in RELAP5 control volume</i>	<i>N</i>
<i>mdbvol(ir5vec(ncon)+filndx(4))=mm</i>	<i>define first index in debcom common block that corresponds with RELAP5 control volume with index of ir5vec(ncon)+filndx(4)</i>	<i>N</i>

**Table 3-2.** Modifications of subroutine EGEN2 for modeling convective and radiative cooling. (continued)

Line of Fortran	Comments	Line status <sup>a</sup>
$ndbvol(ir5vec(ncon)+filndx(4))=nn$	define second index in deb-com common block that corresponds with RELAP5 control volume with index of $ir5vec(ncon) + filndx(4)$	N
$tmpdeb(mm,nn)=tz(i)$	define temperature	N
$porvol(mm,nn)=pore(n)$	define porosity	N
$ddbvol(mm,nn)=dimpe(n)$	define particle size of debris	N
$aovrdb(mm,nn)=(1.-pore(n))*6./dimpe(n)$	define surface area to volume ratio	N
$voldeb(mm,nn)=bv(i)$	define volume of debris	N
$call htrc3b(mm, nn)$	(mm, nn) = indices identifying location of debris in framework of SCDAP data base	N
$voldeb=2\pi *vole(1,n)$	1=i-th node, 2=k-th node, etc.	N
$qcoupl(ir5)=qcoupl(ir5)+qnchdb(mm,nn)$	total heat to RELAP5 volume	N
$qwgcou(ir5)=qwgcou(ir5)+qfgdeb(mm,nn)$	heat to vapor phase	N
$gamcou(ir5)=gamcou(ir5)+gmwdeb(mm,nn)*2\pi *bv(i)/v(numdbv(mm,nn))$	volumetric vapor generation rate, v=volume of RELAP5 control volume	N
$bg(i)=bg(i)-qnchdb(mm,nn)/bv(i)$	adjust power for heat loss, COUPLE nodes map one for one into indices (mm,nn)	N
repeat lines beginning with "ir5=0" and ending with line above for nodes j, k, and l		N
a: E = existing line, M = modified line, N = new line		

Input variables need to be defined for the existing subroutine HTRC3B in order to use this subroutine for calculating the heat transfer between porous debris and fluid in the lower head of the reactor vessel. These input variables need to be added to the common block named debcom. This common block also stores the variables output by subroutine HTRC3B. Table 3-3 lists the input and output variables in this common block that need to be defined or applied by the COUPLE model. The input variables include; (1) porosity of debris, (2) particle size of debris, (3) surface area to volume ratio of debris, (4) volume of debris, (5) temperature of debris, and (6) index of RELAP5 volume within which the debris is located. The

output variables are; (1) heat transfer between the debris and fluid, (2) heat transfer between debris and vapor phase of the fluid, and (3) volumetric vapor generation rate.

**Table 3-3.** Variables in common block debcom that are input and output variables for subroutine HTRC3B for porous debris heat transfer.

Variable name	Units	Category	Variable definition
<i>porvol(m,n)</i>	<i>unitless</i>	<i>input</i>	<i>porosity of debris at location defined by indices (m,n)</i>
<i>ddbvol(m,n)</i>	<i>m</i>	<i>input</i>	<i>diameter of debris particles (m)</i>
<i>aovrdb(m,n)</i>	<i>(1/m)</i>	<i>input</i>	<i>surface area of debris per unit volume of debris</i>
<i>tmpdeb(m,n)</i>	<i>K</i>	<i>input</i>	<i>temperature of debris (K)</i>
<i>voldeb(m,n)</i>	<i>m<sup>3</sup></i>	<i>input</i>	<i>volume of debris</i>
<i>nvoldb(n)</i>	<i>-</i>	<i>input</i>	<i>number of RELAP5 control volumes in stack of control volumes represented by index n</i>
<i>numdbv(m,n)</i>	<i>-</i>	<i>input</i>	<i>index of RELAP5 control volume at location (m,n)</i>
<i>qnchdb(m,n)</i>	<i>W</i>	<i>output</i>	<i>rate of heat transfer between debris and fluid.</i>
<i>qfgdeb(m,n)</i>	<i>W</i>	<i>output</i>	<i>rate of heat transfer between debris and vapor phase of coolant</i>
<i>gmwdeb(m,n)</i>	<i>(kg/m<sup>3</sup>.s)</i>	<i>output</i>	<i>volumetric vapor generation rate</i>

The input and output variables in common block debcom are defined for each SCDAP and COUPLE node with porous debris. The node locations are identified by indices that define the radial and axial position of the debris. For in-core debris, these indices are mapped according to axial nodes and radial segments in the core region. For debris represented by the COUPLE model, these indices are mapped as a function of the COUPLE nodes. The indices used for debris represented by the COUPLE model will not overlap those used for in-core debris. In order to conserve computer memory, the lower bound of the indices for debris represented by the COUPLE model will be contiguous with the upper bound of the indices used to represent in-core debris. As shown previously, Table 3-2 shows the basic structure of fortran programming that will be added to subroutine EGEN2 to determine the indices for the debcom common block variables and define the values of the required debcom variables. Subroutine HTRC3B will be modified to have an additional input variable that defines the starting value of the index m in the do500 do loop in this subroutine. This modification is described in Table 3-4. In the call to subroutine HTRC3B from subroutine SCDAD5, the input argument mstart will be defined to have a value of 1. The basic structure of subroutine HTRC3B is described in Table 3-5.

**Table 3-4.** Fortran modifications to subroutine HTRC3B for porous debris heat transfer.

Line of Fortran	Comments	Status <sup>a</sup>
<i>subroutine htrc3b(mstart, n)</i>	<i>add mstart to argument list, where mstart is starting value of index m</i>	<i>M</i>

**Table 3-4.** Fortran modifications to subroutine HTRC3B for porous debris heat transfer. (continued)

Line of Fortran	Comments	Status <sup>a</sup>
<i>do 500 m=mstart,nvoldb(n)</i>	<i>make starting value of index m a function of input argument instead of always being equal to 1</i>	<i>M</i>
<i>a: M = modified line</i>		

**Table 3-5.** Basic structure of subroutine HTRC3B for calculating debris to fluid heat transfer.

Line of Fortran	Comments
<i>subroutine htrc3b(mstart, n)</i>	<i>mstart = starting value of index m in do500 do loop</i>
<i>*call comctl</i>	<i>access RELAP5 variables</i>
<i>*call contrl</i>	<i>access RELAP5 variables</i>
<i>*call voldat</i>	<i>access RELAP5 variables</i>
<i>*call fast</i>	<i>access RELAP5 variables</i>
<i>*call ufiles</i>	<i>access RELAP5 variables</i>
<i>iv=ir5vc1+filndx(4)</i>	<i>index for RELAP5 variables in voldat common block</i>
<i>block of coding to determine heat transfer regime based on criteria defined in Table 2-1; regime is function of volume fraction of liquid and debris temperature</i>	
<i>block of coding that calculates convective heat transfer for each heat transfer regime according to correlations defined in Section 2,</i>	
<i>block of coding that calculates the heat transferred to liquid and vapor phases by radiation heat transfer</i>	
<i>block of coding that calculates total rate of heat transfer to fluid at the specified location in debris, also total heat transfer to vapor phase of fluid, and volumetric vapor generation rate</i>	
<i>end of subroutine</i>	

The changes made to the argument list of subroutine EGEN2 require that changes be made to the call of subroutine EGEN2 from subroutine COUPLE. These changes are described in Table 3-6.

**Table 3-6.** Fortran changes in subroutine COUPLE for implementing new models for heat transfer in porous debris.

Line of Fortran	Comments	Status <sup>a</sup>
<i>call egen2(a(i8), a(i107), a(i27), a(i28), a(i29), a(n20),</i>	<i>extend call to egen2 to account for variables needed to model heat transfer to fluid</i>	<i>E</i>
<i># a(n103), a(i110), a(iratpo),</i>	<i>variable a(iratpo) is no longer the last argument</i>	<i>M</i>
<i># a(i22), a(i70), a(i72), a(iqcopt), a(iqwcp),</i>	<i>add to call pointers needed for debris to fluid heat transfer</i>	<i>N</i>
<i># a(igmcp), a(i108), a(iqcrdb), a(ivcnpt), a(iptihs), a(iptpor))</i>	<i>"</i>	<i>N</i>
<i>a: E = existing line of fortran, M = modified line, N = new line.</i>		

Several subroutines require minor modifications for implementation of porous debris to fluid heat transfer. Subroutine COUPLE needs to have the call to DBVPGN skipped and the variable qd set to zero when the option to use the advanced porous debris heat transfer model is defined by the code user. In subroutine COUQOT after start of do820 do loop, a statement needs to be added to skip the calculation of qcoup, qwgcou and gamcou when ihpore(n) (indicator of calculation of debris to fluid heat transfer) is equal to 1; when debris to fluid heat transfer has been calculated, then the calculation of convective heat transfer at surface is redundant. In do80 do loop of subroutine CG2, the variable ihpore(i) needs to be set to zero. In the do10 do loop of subroutine ICPL, the variable qcrdeb(i) needs to be set to 0.0. In subroutine RGEN after initialization of variable ipfrto (pointer to multiplier on RELAP5 power density table for each node in COUPLE mesh), the initialization of the variable iqcrdb (pointer to value of heat transfer from debris to fluid) needs to be added in manner parallel to initialization of ipfrto. In subroutine CONSET, the variable iptpor (pointer to array ihpore) needs to be defined in manner parallel to definition of the pointer iptihs (pointer to array ihsave). The common block iparm needs the integer variables iqcrdb and iptpor added to it. The indicator of the option to be used for calculating debris to fluid heat transfer (variable ndbhr) needs to be defined by input in subroutine CONSET in the A.23.8.2 block of data. The subroutine MAJCOU needs to be extended to print for every node with porous debris the ratio of heat transfer to fluid to internal heat generation (qcredeb(i)/bg(i)). For the case of zero power, the printout would not be performed. An input subroutine needs to be extended to indicate whether the heat transfer from fluid to debris should be calculated by the simplified method or the detailed method.

The heat transfer between the liquid and vapor phases of the fluid in porous debris will be modeled in an approximate manner. The thermal equilibrium model will be used. The thermal equilibrium model is

employed by setting the interfacial heat transfer coefficient to a high value. In particular, the fortran lines shown in Table 3-7 are added to subroutine PHANTV.

**Table 3-7.** Fortran lines added to RELAP5 subroutine PHANTV to model interphase heat transfer in porous debris.

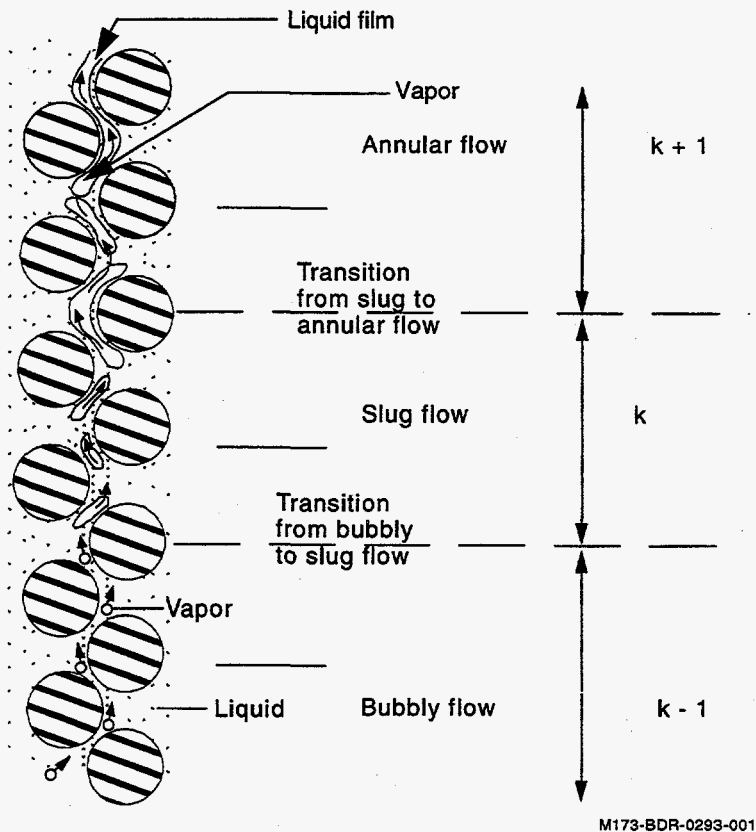
Fortran line	Comments	Status <sup>a</sup>
<i>337 continue</i>		<i>E</i>
<i>endif</i>		<i>E</i>
<i>if(idbvol(i).gt.0)then</i>	<i>after above two lines, check for possibility of porous debris in RELAP5 volume with index i</i>	<i>N</i>
<i>hif(i)=1.0e+9</i>	<i>set interfacial heat transfer coefficient to large value</i>	<i>N</i>
<i>hig(i)=1.0e+9</i>	<i>set interfacial heat transfer coefficient to large value</i>	<i>N</i>
<i>endif</i>		
<i>a: E = existing line, N = new line</i>		

## 4. Flow Losses and Interphase Drag in Porous Debris

This section defines extensions for the modeling of flow losses and interphase drag in porous debris regions in order to more accurately calculate the heatup of the porous debris. The flow losses and interphase drag are a function of the flow regime. First, the algorithm to be used to identify the flow regime for all possible ranges of coolant conditions will be presented. Second, the algorithm used to calculate the flow losses due to contact of the fluid with the debris will be described. Third, the algorithm to be used to calculate interphase drag will be described. These algorithms are presented in abbreviated form in this report; a complete description of these algorithms is presented in Reference 2.

### 4.1 Flow regime identification

The flow regime is assumed to be identified by the volume fraction of vapor in the fluid and by the heat transfer regime. The heat transfer regime is determined by the algorithm presented in Section 2. For pre-CHF heat transfer regimes, five flow regimes are assumed to be possible in porous debris. These five regimes are; (1) bubbly flow, (2) bubbly-slug flow, (3) slug flow, (4) slug-annular flow, and (5) annular flow. A schematic of the flow regimes is shown in Figure 4-1. In the figure, "k" is a node number that identifies the location of porous debris. For post-CHF heat transfer regimes, five flow regimes are also identified; (1) inverted annular flow, (2) transition from inverted annular flow to inverted slug flow, (3) inverted slug flow, (4) inverted slug-mist flow, and (5) mist flow. A schematic of these flow regimes is shown in Figure 4-2.



**Figure 4-1.** Schematic of pre-surface dryout flow regimes.

The pre-CHF flow regimes are distinguished by the volume fraction of vapor as defined by the following equations;

Bubbly Flow

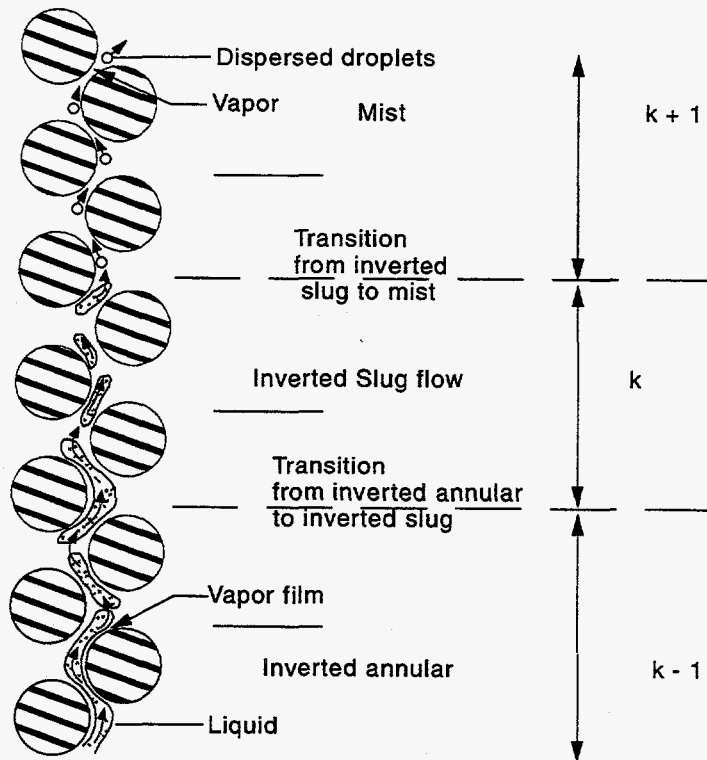
$$0 < \alpha_g < \alpha_1 \tag{4-1}$$

where

$$\alpha_1 = \begin{cases} 0.6(1 - \gamma)^2 & \gamma > 0.29 \\ 0.3 & \gamma \leq 0.29 \end{cases} \text{ and}$$

$$\gamma = D_b / D_p, \text{ and}$$

$$D_b = 1.35 \left[ \frac{\sigma}{g(\rho_f - \rho_g)} \right]^{0.5}$$



M173-BDR-0293-002

**Figure 4-2.** Schematic of post-surface dryout flow regimes.

where

- $D_b$  = bubble diameter (m),
- $D_p$  = particle diameter (m),
- $\alpha_g$  = volume fraction of vapor in fluid,
- $\sigma$  = surface tension ( $\text{kg/s}^2$ ),
- $g$  = gravitational acceleration ( $\text{m/s}^2$ ),
- $\rho_f$  = density of liquid ( $\text{kg/m}^3$ ),
- $\rho_g$  = density of gas ( $\text{kg/m}^3$ ).

Bubbly-Slug Flow

$$\alpha_1 \leq \alpha_g < \alpha_2$$

(4-2)



where

$$\alpha_2 = \pi/6.$$

Slug Flow

$$\alpha_2 \leq \alpha_g < \alpha_3 \tag{4-3}$$

where

$$\alpha_3 = 0.6.$$

Slug-Annular Flow

$$\alpha_3 \leq \alpha_g < \alpha_4 \tag{4-4}$$

where

$$\alpha_4 = 0.925.$$

Annular Flow

$$\alpha_4 \leq \alpha_g \leq 1. \tag{4-5}$$

According to Reference 2, the post-CHF flow regimes are distinguished by the volume fraction of vapor as shown by the following equations;

Inverted Annular Flow

$$0 < \alpha_g < \alpha_1 \tag{4-6}$$

where  $\alpha_1$  is given in Equation (4-1).

Inverted Annular-Inverted Slug Flow

$$\alpha_1 \leq \alpha_g < \alpha_2 \tag{4-7}$$

where  $\alpha_2 = \pi/6$ .

### Inverted Slug Flow

$$\alpha_2 \leq \alpha_g < \alpha_3 \quad (4-8)$$

where  $\alpha_3 = 0.6$ .

### Inverted Slug-Mist Flow

$$\alpha_3 \leq \alpha_g < \alpha_4 \quad (4-9)$$

where  $\alpha_4 = 0.925$ .

### Mist Flow

$$\alpha_4 \leq \alpha_g \leq 1. \quad (4-10)$$

Reference 3 recommends that the post-CHF flow regime be distinguished by three thresholds for void fraction, namely  $\alpha_1$ ,  $\alpha_2$  and  $\alpha_3$ . For  $\alpha_1 \leq \alpha_g < \alpha_2$ , the flow regime is inverted annular flow. For  $\alpha_2 \leq \alpha_g < \alpha_3$ , the flow regime is inverted slug flow. For  $\alpha_g > \alpha_3$ , the flow regime is mist flow. The value of  $\alpha_1$  is approximately given by the equation

$$\alpha_1 = \frac{(1 - \varepsilon)3\delta}{\varepsilon D_p} \quad (4-11)$$

where (4-12)

- $\alpha_1$  = threshold void fraction for inverted annular flow,
- $\varepsilon$  = porosity of debris,
- $\delta$  = vapor film thickness (m),
- $D_p$  = size of debris particles (m).

Assuming that the liquid field does not contain vapor bubbles and that the debris bed consists of packed spherical particles, the vapor film thickness is related to the volume fraction of vapor by the equation

$$[0.5D_p + \delta]^3 = 0.125D_p^3 (0.91\alpha_g + 1)$$

The values of  $\alpha_2$  and  $\alpha_3$  need to be determined empirically. A suggested range for  $\alpha_3$  is 0.9 - 0.95.

If the latter method for calculating post-CHF flow regimes can be successfully implemented, then the former method does not need to be implemented.

## 4.2 Models for Flow Losses

The resistance applied to the flow of liquid and vapor phases of the fluid due to contact with the debris and with each other is a function of the flow regime and velocities of the liquid and vapor phases.<sup>4,5,11</sup> The flow losses are calculated in terms of drag force (pressure loss gradient) in the porous debris. In the equations below, this drag force is represented by the terms  $F_{pg}$  and  $F_{pl}$  for the vapor and liquid phases of the fluid, respectively. These terms have the units of  $N/m^3$ .

### 4.2.1 Drag Force for Superheated Steam (Single-Phase).

The gas-phase drag force can be calculated from the Kozeny-Carman equation,<sup>4</sup> which can be written as

$$F_{pg} = \varepsilon [a \mu_g j_g + b \rho_g j_g^2] \quad (4-13)$$

where the constants a and b are defined as

$$a = 150 \frac{(1 - \varepsilon)^2}{\varepsilon^3 D_p^2} \quad \text{and} \quad b = 1.75 \frac{(1 - \varepsilon)}{\varepsilon^3 D_p}$$

- $F_{pg}$  = flow resistance ( $N/m^3$ ),
- $j_g$  = superficial velocity of the gas (m/s),
- $\rho_g$  = density of the gas ( $kg/m^3$ ),
- $\mu_g$  = viscosity of the gas ( $kg/m \cdot s$ ).

The superficial velocity  $j_g$  is related to the  $v_g$  (velocity of steam in the interstices) by the equation;

$$j_g = \varepsilon v_g. \quad (4-14)$$

### 4.2.2 Drag Force for Subcooled and Saturated Liquid (Single-Phase)

The debris drag force acting on the single-phase is calculated by the equation

$$F_{pf} = \varepsilon[a\mu_f j_f + b\rho_f j_f |j_f|] \quad (4-15)$$

where

$$\begin{aligned} j_f &= \text{superficial velocity of the liquid (m/s),} \\ \rho_f &= \text{density of the liquid (kg/m}^3\text{),} \\ \mu_f &= \text{viscosity of the liquid (kg/m} \cdot \text{s).} \end{aligned}$$

### 4.2.3 Drag Force for Two-Phase Flow

The friction between the two phases and the debris can be modeled with two distinct drag components<sup>4</sup>, i.e., particle-gas, and particle-liquid. The particle-gas drag force ( $F_{pg}$ ) is the drag force by the particles on the gas through the liquid layer, which is opposed by an equal and opposite force applied by the particles on the other side of the liquid layer. The particle-liquid drag force ( $F_{pl}$ ) is the force by particles on the liquid due to liquid motion, i.e., the liquid drag force against the particles due to liquid motion.

#### 4.2.3.1 Particle-Gas Drag Force for Two-Phase Flow

The Kozeny-Carman equation for the particle-gas drag force per unit of total debris bed volume can be written as

$$F_{pg} = \varepsilon \left[ \frac{a\mu_g j_g}{k_g} + \frac{b\rho_g j_g^2}{\eta_g} \right] \quad (4-16)$$

where

$$a = \frac{150(1-\varepsilon)^2}{\varepsilon^3 D_p^2},$$

$$b = \frac{1.75(1-\varepsilon)}{\varepsilon^3 D_p}.$$

The variables  $k_g$  and  $\eta_g$  are the relative permeabilities, which vary with different flow regimes.

The relative permeabilities may be explicitly written as follows.

#### For Bubbly and Slug Flows

$$k_g = \left( \frac{1-\varepsilon}{1-\varepsilon\alpha_g} \right)^{4/3} \alpha_g^3 \quad \text{and} \quad \eta_g = \left( \frac{1-\varepsilon}{1-\varepsilon\alpha_g} \right)^{2/3} \alpha_g^3. \quad (4-17)$$

#### For Slug-Annular Flow

$$k_g = \frac{\left(\left(\frac{1-\varepsilon}{1-\varepsilon\alpha_g}\right)^{4/3}\right)\alpha_g^2}{\left(W + \frac{1-W}{\alpha_g}\right)} \text{ and } \eta_g = \frac{\left(\frac{1-\varepsilon}{1-\varepsilon\alpha_g}\right)^{2/3}\alpha_g^2}{\left(W + \frac{1-W}{\alpha_g}\right)} \quad (4-18)$$

where W is the weighting function written as

$$W = \xi^2(3 - 2\xi), \text{ and } \xi = \frac{\alpha_g - \alpha_3}{\alpha_4 - \alpha_3}.$$

#### For Pure Annular Flow

$$k_g = \left(\frac{1-\varepsilon}{1-\varepsilon\alpha_g}\right)^{4/3}\alpha_g^2 \text{ and } \eta_g = \left(\frac{1-\varepsilon}{1-\varepsilon\alpha_g}\right)^{2/3}\alpha_g^2. \quad (4-19)$$

#### For Inverted Annular Flow, Inverted Annular-Inverted Slug Flow, Inverted Slug Flow, and Mist Flow

The particle-gas drag relationships for these surface-dryout regimes will be treated in a similar fashion to the corresponding presurface-dryout regimes except that the roles of vapor and liquid are interchanged.

#### **4.2.3.2 Particle-Liquid Drag Force for Two-Phase Flow**

The equation for the particle-liquid drag force per unit of total debris bed volume can be written as

$$F_{pf} = \varepsilon \left[ \frac{a\mu_f j_f}{k_f} + \frac{b\rho_f j_f |j_f|}{\eta_f} \right] \quad (4-20)$$

The variables  $k_f$  and  $\eta_f$  are the relative permeabilities, which vary with different flow regimes.

#### For Bubbly, Bubbly-Slug, Slug, Slug-Annular, and Annular Flow

A single expression for the relative permeabilities for these regimes can be written as

$$k_f = \eta_f = (1 - \alpha_g)^3. \quad (4-21)$$

#### For Inverted Annular Flow

$$k_f = \left( \frac{1-\varepsilon}{1-\varepsilon\alpha_f} \right)^{4/3} \alpha_f^2 \text{ and } \eta_f = \left( \frac{1-\varepsilon}{1-\varepsilon\alpha_f} \right)^{2/3} \alpha_f^2 \quad (4-22)$$

where

$\alpha_f$  = volume fraction of liquid phase.

#### For Inverted Annular-Inverted Slug Flow

$$k_f = \frac{\left( \frac{1-\varepsilon}{1-\varepsilon\alpha_f} \right)^{4/3} \alpha_f^2}{\left( W + \frac{1-W}{\alpha_g} \right)} \text{ and } \eta_f = \frac{\left( \frac{1-\varepsilon}{1-\varepsilon\alpha_f} \right)^{2/3} \alpha_f^2}{\left( W + \frac{1-W}{\alpha_f} \right)} \quad (4-23)$$

where W is the weighting function written as

$$W = \xi^2(3 - 2\xi) \text{ and } \xi = \frac{\alpha_f - \alpha_3}{\alpha_4 - \alpha_3}$$

#### For Inverted Slug and Mist Flow

$$k_f = \left( \frac{1-\varepsilon}{1-\varepsilon\alpha_f} \right)^{4/3} \alpha_f^3 \text{ and } \eta_f = \left( \frac{1-\varepsilon}{1-\varepsilon\alpha_f} \right)^{2/3} \alpha_f^3 \quad (4-24)$$

### 4.3 Models for Interphase Drag

The liquid-gas interfacial drag force ( $F_i$ ) is the force by the liquid on the gas due to relative motion between the two-phases.

#### For Bubbly Flow

For the bubbly liquid-gas interfacial drag force, the drag force exerted on a single bubble is multiplied by the number of bubbles per unit volume of the porous layer.

$$F_i = C_v \frac{\mu_f j_s}{D_b^2} + C_i \frac{(\rho_f - \alpha_g \rho_f + \alpha_g \rho_g) j_s^2}{D_b^2 \varepsilon} \quad (4-25)$$

where  $j_s$  is the drift velocity of the bubble relative to the mixture written as

$$j_s = j_g \frac{(1 - \alpha_g)}{\alpha_g} - j_f$$

and the coefficients  $C'_v$  and  $C'_i$  are expressed as

$$\left. \begin{aligned} C'_v &= 18\alpha_g f \\ C'_i &= 0.34(1 - \alpha_g)^3 \alpha_g f^2 \end{aligned} \right\} \text{ for } 0 < \alpha_g \leq \alpha_0$$

$$\left. \begin{aligned} C'_v &= 18(\alpha_0 f + \alpha_g - \alpha_0) \\ C'_i &= 0.34(1 - \alpha_g)^3 (\alpha_0 f^2 + \alpha_g - \alpha_0) \end{aligned} \right\} \text{ for } \alpha_0 \leq \alpha_g \leq \alpha_1$$

where  $\alpha_1$  is defined by Equation (4-1) and  $\alpha_0$  is the void fraction corresponding to the maximum number of bubbles supported by the surface of the particles. This can be expressed as

$$\alpha_0 = \frac{\pi(1 - \varepsilon)}{3\varepsilon} \gamma(1 + \gamma)[6\eta - 5(1 + \gamma)] \text{ for } \alpha_0 \geq 0.$$

In the above equation,

$$\gamma = D_b/D_p, \text{ and } \eta = \left[ \frac{\pi\sqrt{2}}{6(1 - \varepsilon)} \right]^{1/3}.$$

The geometric factor,  $f$ , is given by

$$f = \frac{1}{2}(1 + \gamma) \ln \left( 1 + \frac{2}{\gamma} \right). \quad (4-26)$$

#### For Bubbly-Slug Flow

Slugs are considered as long, thin ellipsoids whose lateral dimension is  $D_b$  and length is  $L_b (=8D_b)$ . The drag force acting on the bubbly-slug flow regime can be expressed by Equation (4-25), with a smooth transition between bubbly and slug flow, by using a weighting function for the coefficients. The coefficients are modified as follows.

$$\left. \begin{aligned} C'_v &= 18(\alpha_0 f + \alpha_g - \alpha_0)(1 - W) + 5.21\alpha_g W \\ C'_i &= (1 - \alpha_g)^3 \{0.34(\alpha_0 f^2 + \alpha_g - \alpha_0)(1 - W) + 0.92\alpha_g W\} \end{aligned} \right\} \quad (4-27)$$

where

$$W = \xi^2(3 - 2\xi) \text{ and } \xi = \frac{\alpha_g - \alpha_1}{\alpha_2 - \alpha_1}.$$

#### For Slug Flow

Since the slugs are fairly long and extend beyond a pore length, they do not flow along the particles as do the spherical bubbles at void fractions less than  $\alpha_0$ . Thus, a geometrical correction on  $j_s$  is not

needed. The drag force can be expressed by Equation (4-25) with modified coefficients  $C'_v$ ,  $C'_i$ , which can be written as

$$C'_v = 5.21\alpha_g \text{ and } C'_i = 0.92(1 - \alpha_g)^3 \alpha_g.$$

#### For Slug-Annular Flow

For the transition from slug to annular flow regime, the weighting function defined for particle-gas drag force can be written as

$$F_i = \left[ 5.21\alpha_g \frac{\mu_f(1-W)}{D_b^2} + \frac{\epsilon a \mu_g W}{k_g} \right] j_s \quad (4-28)$$

$$+ \left[ 0.92\alpha_g(1 - \alpha_g)^3 \frac{(\rho_f - \rho_f \alpha_g + \rho_g \alpha_g)(1-W)}{\epsilon D_b} + \frac{\epsilon \alpha_g b \rho_g W}{1 - \alpha_g \eta_g} \right] j_s^2$$

where  $k_g$ ,  $\eta_g$  and  $W$  are defined as for Equation (4-18) and "a" is defined as for Equation (4-16).

#### For Annular Flow

In annular flow, the reference velocity must be modified to account for the slip between the two phases.

$$F_i = \frac{\epsilon a \mu_g}{k_g} j_s + \frac{\epsilon \alpha_g}{1 - \alpha_g} \left( \frac{b \rho_g}{\eta_g} \right) j_s^2 \quad (4-29)$$

where  $k_g$  and  $\eta_g$  are given by Equation (4-19).

#### For Inverted Annular Flow, Inverted Annular-Inverted Slug Flow, Inverted Slug Flow, and Mist Flow

The interface drag relationships for these post-dryout regimes will be treated in a similar fashion to the corresponding pre-dryout regimes except that the roles of vapor and liquid are interchanged.

## 5. Implementation of Models for Flow Losses and Interphase Drag

The momentum losses and gains to the fluid flowing through porous debris will be represented by the HLOSSF and HLOSSG terms<sup>5</sup> in the RELAP5 field equations. These terms are designed to account for momentum losses due to expansions or contractions of flow areas. The momentum losses of fluid flowing through porous debris in the core region will be taken into account by these terms.



The RELAP5 term for HLOSSG is calculated by the equation;

$$\overset{H}{\Delta} \text{LOSSG} = 0.5K_{Lg}v_g \quad (5-1)$$

where

HLOSSG = form or frictional losses (m/s),

$v_g$  = velocity of the vapor phase (m/s),

$K_{Lg}$  = loss coefficient corresponding with velocity  $v_g$  (unitless).

The RELAP5 term HLOSSF is calculated in a parallel manner.

## 5.1 Single Phase

For single phase gas, the loss coefficient is defined from the Bernoulli equation;

$$\varepsilon \frac{\Delta P}{\Delta x} + 0.5\varepsilon K_{Lg} \alpha_g \rho_g v_g^2 / \Delta x = 0 \quad (5-2)$$

where

$\Delta P$  = pressure drop as fluid flows the distance  $\Delta x$  at velocity of  $v_g$  (N/m<sup>2</sup>),

$\Delta x$  = distance (m)

The term  $\varepsilon(\Delta P/\Delta x)$  in the above equation is equal to the term  $-F_{pg}$  in Section 4.2. Thus,

$$F_{pg} = 0.5\varepsilon K_{Lg} \alpha_g \rho_g v_g^2 / \Delta x. \quad (5-3)$$

Solving the above equation for  $K_{Lg}$ , the result is

$$K_{Lg} = \frac{2\Delta x F_{pg}}{\varepsilon \alpha_g \rho_g v_g^2}. \quad (5-4)$$

The loss coefficients for the vapor phase are stored in the RELAP5 variable formgj(i), where "i" is the junction index. The variable formgj(i) is calculated in subroutine HLOSS. For junctions that are

connected with control volumes containing porous debris, fortran coding will be added to subroutine HLOSS that calculates formgj(i) by the equation

$$\text{formgj}(i) = K_{Lg} = \frac{2\Delta x F_{pg}}{\epsilon \alpha_g \rho_g v_g^2} \quad (5-5)$$

The term  $F_{pg}$  is from Section 4.2. For the single phase case,  $\alpha_g = 1$ . In calculating formgj(i),  $\Delta x$  is the distance of flow through porous debris represented by junction "i", and  $\rho_g$  and  $v_g$  are the vapor density and vapor velocity at the junction, respectively.

The loss coefficients for single phase liquid are calculated in a manner parallel to that for single phase vapor. They are stored in the RELAP5 variable formfj(i), which is calculated by the equation

$$\text{formfj}(i) = K_{Lf} = \frac{2\Delta x F_{pf}}{\epsilon \alpha_f \rho_f v_f^2} \quad (5-6)$$

This implementation of  $F_{pg}$  and  $F_{pf}$  into the RELAP5 momentum equations is consistent with the current implementation of other such losses, i.e., a partially implicit method is used where  $v_g^2$  is implemented as  $v_g^n v_g^{n+1}$  and  $v_f^2$  is implemented as  $v_f^n v_f^{n+1}$ .

## 5.2 Two Phase

The steady state phasic porous debris momentum equations (assume horizontal flow, no gravity) are of the form

$$\epsilon \alpha_g \frac{\Delta P}{\Delta X} + F_{pg} + F_i = 0 \quad (5-7)$$

$$\epsilon \alpha_f \frac{\Delta P}{\Delta X} + F_{pf} - F_i = 0 \quad (5-8)$$

where

$F_{pg}$ ,  $F_{pf}$ , and  $F_i$  are from Section 4.2.

For the situation of two-phase flow in porous debris, the RELAP5 variables formfj(i) and formgj(i) will be calculated in a modified manner to approximate interphase drag. For this situation, formfj(i) and formgj(i) will be calculated by the equations

$$\text{formgj}(i) = \frac{2\Delta x(F_{pg} + F_i)}{\epsilon \alpha_g \rho_g v_g^2} \quad (5-9)$$

$$\text{formfj}(i) = \frac{2\Delta x(F_{pf} - F_i)}{\epsilon \alpha_f \rho_f v_f^2} \quad (5-10)$$

where

$F_i$  = interphase drag ( $N/m^3$ ).

The terms  $F_{pg}$ ,  $F_{pf}$  and  $F_i$  are from Section 4.2.

As with the single phase case, this implementation of  $F_{pg}$  and  $F_{pf}$  into the RELAP5 is consistent with the current implementation of other such losses, i.e., a partially implicit method is used where  $v_g^2$  is implemented as  $v_g^n v_g^{n+1}$  and  $v_f^2$  is implemented as  $v_f^n v_f^{n+1}$ .

Using this implementation [Equation(5-9) and (5-10)], the porous debris interphase drag term is not as consistent as the current interphase drag term in RELAP5. For this approximate implementation it is assumed that the phase velocities change very little between time steps (slow transients) i.e.,  $v_g^{n+1} \approx v_g^n$  and  $v_f^{n+1} \approx v_f^n$ . For fast transients, where the phasic velocities change significantly between time steps, the implementation of the porous debris interphase drag needs to be consistent with the current interphase drag implementation in RELAP5, i.e.,  $(v_g - v_f)^2$  is implemented as  $(v_g - v_f)^n (v_g - v_f)^{n+1}$ , where  $v_g^{n+1} \neq v_g^n$  and  $v_f^{n+1} \neq v_f^n$ . This implementation will need to be done in subroutine phantj rather than subroutine hloss. This is a task for future work. In the interim, a smaller time step will be applied during periods of rapidly changing velocities.

A branch will be put into the first part of the do 2000 loop in subroutine HLOSS to calculate  $\text{formfj}(i)$  and  $\text{formgj}(i)$  for RELAP5 junctions that represent flow through porous debris regions. The basic structure of the Fortran changes required for subroutine HLOSS are shown in Table 5-1. The programming subset in subroutine FWDRAG to identify the flow regime for a porous debris region ( $\text{idbvol}(i)=1$ ) will be removed from subroutine FWDRAG and implemented into subroutine HLOSS.

**Table 5-1.** Modification of subroutine HLOSS for modeling of flow losses in porous debris.

Fortran coding	Comments	Line status <sup>a</sup>
<i>*call voldat</i>	RELAP5 volume variable	E
<i>*call scddat</i>	after above line, add common block storing length of arrays storing debris indices and characteristics	N
<i>*call debcom</i>	add common block storing debris indices and characteristics	N

**Table 5-1.** Modification of subroutine HLOSS for modeling of flow losses in porous debris. (continued)

Fortran coding	Comments	Line status <sup>a</sup>
<i>*call tblsp</i>	<i>add common block storing debris indices and characteristics</i>	<i>N</i>
<i>do 2000 m = 1,lvptr(il)</i>	<i>start of do2000 loop</i>	<i>E</i>
<i>.....</i>	<i>skip display of some lines</i>	<i>E</i>
<i>kx = k + iand(ishft(jcex(i),-13),3)</i>	<i>kx = index of RELAP5 volume connected to junction i, after adjusting for flow direction, kx is replaced with ky</i>	<i>E</i>
<i>lx = l + iand(ishft(jcex(i),-10),3)</i>	<i>lx = index of another RELAP5 volume connected to junction i, after adjusting for flow direction, lx is replaced with ly</i>	<i>E</i>
<i>if (vf.lt. 0.0) then</i>	<i>beginning of "if()then" block</i>	<i>E</i>
<i>...</i>	<i>contents of "if()then" block</i>	<i>E</i>
<i>endif</i>	<i>end of "if()then" block</i>	<i>E</i>
<i>if(idbvol(ky).eq.2)then</i>	<i>check to see whether junction connected with volume containing porous debris, idbvol(ky)=2=yes, place this and following lines immediately after above "if()then" block</i>	<i>N</i>
<i>epsdb=porvol(mdbvol(ky),ndbvol(ky))</i>	<i>epsdb = porosity of debris</i>	<i>N</i>
<i>diadb=ddbvol(mdbvol(ky),ndbvol(ky))</i>	<i>diadb = diameter of debris particles (m)</i>	<i>N</i>
<i>velgdb=abs(velgj(i))</i>	<i>velgj(i) = velocity of vapor phase at junction with index i (m/s)</i>	<i>N</i>
<i>if(velgdb.lt.1.0)velgdb=1.0</i>	<i>set lower limit of velocity to lower limit for flow drag correlations</i>	<i>N</i>
<i>velfdb=abs(velfj(i))</i>	<i>velfj(i) = velocity of liquid phase at junction with index i (m/s)</i>	<i>N</i>
<i>if(velfdb.lt.1.0)velfdb=1.0</i>	<i>set lower limit of velocity to lower limit for flow drag correlations</i>	<i>N</i>
<i>fpg=f1(epsdb,diadb,velgdb,voidf(ky),rhog(ky))</i>	<i>fpg=drag force exerted by particles on vapor phase (N/m<sup>3</sup>), calculated using correlations described in Section 4.</i>	<i>N</i>

**Table 5-1.** Modification of subroutine HLOSS for modeling of flow losses in porous debris. (continued)

Fortran coding	Comments	Line status <sup>a</sup>
$f_i=f_2(\text{epsdb}, \text{diadb}, \text{velgdb}, \text{voidf}(ky), \text{rhog}(ky))$	$f_i$ =drag force exerted by liquid phase on vapor, calculated using correlations described in Section 4.	N
$\text{formgj}(i)=\text{formgj}(i)+dl(ky)*(fpg+f_i)/(\text{epsdb}*\text{voidg}(ky)*\text{rhog}(ky)*\text{velgdb}**2)$	calculate form loss for vapor phase for junction $i$ accounting for presence of porous debris (unitless)	N
$f_{pf}=f_3(\text{epsdb}, \text{diadb}, \text{velfdb}, \text{voidf}(ky), \text{rhog}(ky))$	$f_{pf}$ =drag force exerted by particles on liquid phase ( $N/m^3$ ), calculated using correlations described in Section 4	N
$\text{formfj}(i)=\text{formfj}(i)+dl(ky)*(f_{pf}-f_i)/(\text{epsdb}*\text{voidf}(ky)*\text{rhof}(ky)*\text{velfdb}**2)$	calculate form loss for liquid phase for junction $i$ accounting for presence of porous debris (unitless)	N
end if	end of "if(idbvol(ky).eq.2)then" block	N
...	repeat for connecting volume with index $ly$ the sequence of coding beginning with line "if(idbvol(ky).eq.2)then" and ending with line " $\text{formfj}(i)=\text{formfj}(i)+dl(ky)*(f_{pf}-f_i)/(\text{epsdb}*\text{voidf}(ky)*\text{rhof}(ky)*\text{velfdb}**2)$ "	N
if(idbvol(ky).eq.2.or.idbvol(ly).eq.2)then	limit $\text{formgj}(i)$ and $\text{formfj}(i)$ to upper bound values for porous debris and then go to end of do loop	N
if(formgj(i).gt.5000.)formgj(i)=5000.	limit to upper bound value	N
if(formfj(i).gt.5000.)formfj(i)=5000.	limit to upper bound value	N
go to 2000	skip over coding in do loop for volumes without porous debris	N
end if	end if	
a: E = existing line, M = modified line, N = new line		

Since the variables  $\text{formfj}(i)$  and  $\text{formgj}(i)$  account for both interphase drag and wall friction, the RELAP5 variable for interphase drag and the RELAP5 variables for wall friction will be set to zero. The RELAP5 variable in the field equations for interphase drag is named FI and the RELAP5 variable in the field equations for wall friction for the liquid and vapor phases are named FWF and FWG, respectively. To preclude a redundant calculation of wall friction, the RELAP5 fortran variables  $\text{fwalfj}(i)$  and  $\text{fwalgj}(i)$  will be set to zero in subroutine FWDRAG for junctions connected with control volumes that contain porous debris. To preclude a redundant calculation of interphase drag, the RELAP5 fortran variable  $\text{fij}(i)$  will be set to a small value subroutine VEXPLT for junctions connected with control volumes that contain porous debris.

## 6. Extensions to In-Core Porous Debris

The models implemented for calculating the cooling of porous debris in the lower head are also applicable for calculating the cooling of debris in the core region. Only a moderate amount of extra effort is required to implement these models for in-core debris. A few lines in subroutine SCDAD5 need to be changed as indicated in Table 6-1. The calculation of flow losses for in-core porous debris is currently calculated by inactive coding in subroutine FWDRAG. This coding will be deleted and the form loss coefficients for both in-core and lower head debris calculated in subroutine HLOSS.

**Table 6-1.** Modifications of subroutine SCDAD5 for application of detailed models for cooling of porous debris

line of fortran	comments	status <sup>a</sup>
<i>\$if -def,debth</i>	<i>line to be removed</i>	<i>R</i>
<i>idbvol(1) = 2</i>	<i>previous line had "idbvol(1) = 1"</i>	<i>M</i>
<i>mstart=1</i>	<i>place just before line shown below</i>	<i>N</i>
<i>call htrc3b(mstart,n)</i>	<i>previous line had "call htrc3b(n)"</i>	<i>M</i>
<i>a: E= existing line, N = new line, M = modified line, R = line to be removed</i>		

## 7. Assessment of Implemented Models

The new models for cooling of porous debris will be assessed by analysis of two idealized problems, two experiments, and an analysis of a full-plant severe accident. The test matrix for assessment is defined in Table 7-1. The first problem involves a calculation of the heatup and pressure drop of superheated steam flowing through a uniform porous debris bed. The SCDAP/RELAP5 results will be compared with hand calculations of the heatup and pressure drop of the superheated steam. The second problem involves the calculation of boil-off and two-phase flow through a porous debris bed. The SCDAP/RELAP5 method of solution will be evaluated for conservation of energy and for robustness. The third test problem will compare the SCDAP/RELAP5 calculated pressure drop for the case of forced flow of two-phase coolant with that calculated previously by an independent solution.<sup>12</sup> This test problem will perform a steady state analysis of a 1 m deep debris bed with internal heat generation, particle size, and porosity representative of a debris bed resulting from a severe accident in a LWR. The fourth test problem will analyze the heat transfer and flow distribution in a two-dimensional debris bed.<sup>13,14</sup> This test problem performs steady state analyses of a debris bed with particles composed of UO<sub>2</sub> for a range of particle sizes and debris bed heights. The maximum temperature calculated in the debris bed for each case will be compared with that calculated previously by an independent solution.<sup>13,14</sup> The fifth and sixth test problems will compare code calculations with the results of two experiments on porous debris beds with transient two-phase cooling. One of these experiments, designated the BNL debris experiment,<sup>15</sup> involved the quenching of a hot debris bed with a uniform composition and particle size. The sixth test problem, designated the UCLA debris experiment,<sup>16</sup> involved the quenching of a hot debris with a nonuniform composition and particle size. The

The seventh test problem will evaluate the effect of the new models on the behavior calculated for a severe accident in a full-plant.

**Table 7-1.** Matrix of test problems for assessing models for cooling of porous debris.

Problem no.	Problem name	Problem description	Focus of assessment
1	<i>Debris test #1</i>	<i>steam flow through uniform porous debris bed with internal heat generation</i>	<i>compare code calculated pressure drop and rate of heat transfer with results from hand calculations</i>
2	<i>Debris Test #2</i>	<i>boil off of water in deep porous debris bed with uniform heat generation</i>	<i>evaluate degree of conservation of mass and energy, evaluate robustness of calculations for the wide range of two-phase flow conditions that exist in deep debris bed with boil off</i>
3	<i>Debris Test #3</i>	<i>forced flow of two-phase coolant through one-dimensional debris bed</i>	<i>compare SCDAP/RELAP5 calculated pressure drop with that calculated by independent solution confirmed by experimental results</i>
4	<i>Debris Test #4</i>	<i>two-dimensional cylindrical debris bed with internal heat generation immersed in pool of water</i>	<i>compare SCDAP/RELAP5 calculated maximum temperature in debris bed with that calculated by independent solution confirmed by experimental results</i>
5	<i>BNL debris</i>	<i>quenching of hot, uniform debris bed</i>	<i>compare calculated and measured transient temperature distribution in test bundle</i>
6	<i>UCLA debris</i>	<i>quenching of hot, nonuniform debris bed</i>	<i>compare calculated and measured transient temperature distribution in debris bed and transient steam flow rate at top of debris bed</i>
7	<i>Surry TMLB'</i>	<i>TMLB' severe accident in PWR</i>	<i>compare calculation of damage progression with new debris models with that without new debris models</i>

## 8. Summary

Designs were described for implementing models for calculating the heat transfer and flow losses in porous debris. The models can be applied to porous debris in the core region and in the lower head of a reactor vessel. The COUPLE model in SCDAP/RELAP5 represents both the porous and nonporous debris that results from core material slumping into the lower head. Currently, the COUPLE model has the capability to model convective and radiative heat transfer from the surfaces of nonporous debris in a detailed manner and to model only in a simplistic manner the heat transfer from porous debris. In order to advance beyond this simplistic modeling for porous debris, designs were developed for a detailed

calculation of the heat transfer from porous debris to fluid. Correlations were identified for convective heat transfer for the following modes of heat transfer; (1) forced convection to liquid, (2) forced convection to gas, (3) nucleate boiling, (4) transition boiling, and (5) film boiling. Interphase heat transfer is modeled in an approximate manner. A design was also described for implementing a model of heat transfer by radiation from debris to the interstitial fluid. A design was described for implementation of models for flow loss and interphase drag in porous debris. Since the models for heat transfer and flow losses in porous debris in the lower head were designed for general application, a design was also described for implementation of these models to the analysis of porous debris in the core region. A test matrix was proposed for assessing the capability of the implemented models to calculate the heat transfer and flow loss in porous debris. The implementation of the models described in this report is expected to improve the COUPLE code calculation of the temperature distribution in porous debris and in the lower head that supports the debris. The implementation of these models is also expected to improve the calculation of the temperature and flow distribution in porous debris in the core region.

## 9. References

1. The SCDAP/RELAP5 Development Team, "SCDAP/RELAP5/MOD3.2 Code Manual, Volume II: Damage Progression Model Theory," NUREG/CR-6150, Vol. 2, Rev. 1 (INEL-96/0422), July 1998.
2. Seungho Paik and L. J. Siefken, "Extensions to SCDAP/RELAP5 Code for the Modeling of Thermal-Hydraulic Behavior in Porous Debris Beds - Preliminary Design Report," EGG-RAAM-10683, March 1993.
3. V. K. Dhir, "Review of Extensions to SCDAP/RELAP5 Code for the Modeling of Thermal-Hydraulic Behavior in Porous Debris", U. S. Nuclear Regulatory Commission Contract NRC 04 92-045, Task 3, June 1993.
4. V. X. Tung, "Hydrodynamic and Thermal Aspects of Two-Phase Flow Through Porous Media," Ph. D. Thesis, University of California, Los Angeles, 1988.
5. The RELAP5 Development Team, "RELAP5/MOD3 Code Manual, Vol. IV: Models and Correlations , "NUREG/CR-5535, INEL-95/0174, August 1995.
6. D. J. Gunn, "Transfer of Heat or Mass to Particles in Fixed and Fluidized Beds," Int. J. Heat and Mass Transfer, Vol. 21, pp. 467-476, 1978.
7. W. M. Rohsenow, "A Method for Correlating Heat Transfer Data for Surface Boiling of Liquids," Trans. ASME, p. 969, 1952.
8. J. S. Ded and J. H. Lienhard, "The Peak Pool Boiling Heat Flux from a Sphere," AIChE Journal, Vol. 18, No. 2, 1972.
9. V. K. Dhir and J. H. Lienhard, "Laminar Film Condensation on Plane and Axisymmetric Bodies in Non-Uniform Gravity," J. Heat Transfer, Vol. 43, No. 1, 1971.
10. W. Chu, V. K. Dhir, and J. S. Marshall, "Study of Pressure Drop, Void Fraction and Relative Permeabilities of Two-Phase Flow through Porous Media," AIChE Symposium Series, Vol. 79, No. 225, pp. 224-235, 1983.
11. V. X. Tung and V. K. Dhir, "A Hydrodynamic Model for Two-Phase Flow Through Porous Media," Int. J. Multiphase Flow, Vol. 14, No. 1, pp. 47-65, 1988.
12. V. X. Tung, V. K. Dhir, and D. Squarer, "Forced Flow Cooling Studies of Volumetrically Heated Porous Layers," Second International Topical Meeting on Nuclear Reactor Thermal-Hydraulics, Santa Barbara, California, USA, January 11-14, 1983.



13. Mo Chung and Ivan Catton, "Post-Dryout Heat Transfer in a Multi-Dimensional Porous Bed," *Nuclear Engineering and Design*, 128, 1991, pp. 289-304.
14. Ivan Catton and Mo Chung, "Two-Phase Flow in Porous Media with Phase Change: Post-Dryout Heat Transfer and Steam Injection," *Nuclear Engineering and Design*, 151, 1994, pp. 185-202.
15. N. K. Tutu et al., "Debris Bed Quenching under Bottom Flood Conditions (In-Vessel Degraded Core Cooling Phenomenology)," NUREG/CR-3850, 1984.
16. C. H. Wang and V. K. Dhir, "An Experimental Investigation of Multidimensional Quenching of a Simulated Core Debris Bed," *Nuclear Engineering and Design*, 110, 1988, pp. 61-72.