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A UNIFIED MODEL FOR MULTI-PHASE FLOW AND HEAT TRANSFER IN WELLBORES

by

Mohammad Mahbubul Ameen Bachelor of Science, Bangladesh University of Engineering & Technology, 1988.

A Thesis

Submitted to the Graduate Faculty

of the

University of North Dakota

in partial fulfillment of the requirements

for the degree of

Master of Science

Grand Forks, North Dakota

May, 1993

This thesis, submitted by Mohammad Mahbubul Ameen in partial fulfillment of the requirements for the degree of Master of Science from the University of North Dakota, has been read by the Faculty Advisory Committee under whom the work has been done and is hereby approved.

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R. Haran

(Chairperson)

Varlas FMAN

This thesis meets the standards for appearance, conforms to the style and format requirements of the Graduate School of the University of North Dakota, and is hereby approved.

Harney Knu

Dean of the Graduate School 4-28-93

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ABSTRACT

Multiphase flow is a common occurrence in the chemical and petroleum industries. The objective of this study was to apply the principles of multiphase flow to the production of petroleum fluids. A unified model was developed to predict the pressure profiles in wellbores using models available in the literature, which was then used to develop a simulator. A rigorous approach was also taken to model heat transfer and predict the temperature profiles in wellbores under various circumstances.

Our model is capable of predicting the pressure profiles for various channel orientation and geometries. It can handle flow in vertical, and inclined system. Countercurrent flow and flow in downward direction can also be simulated. With appropriate value for the parameters, the model applies to liquid-liquid systems in addition to the gas-liquid systems.

The temperature profile in a wellbore is important to the petroleum industry. Fluid temperature determines various properties such as viscosity, density, the extent of dissolved gases etc. The pressure profile depends on these physical properties. In addition, the temperature profile is important in many production operations in arctic regions. A prior knowledge of the temperature and pressure profile enables the operators to take preventive measures against the clogging of pipelines due to hydrate or wax

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formation. Accurate temperature estimation is also important during such operations as drilling, cementing etc.

Fluid temperature depends on the extent of heat loss from the wellbore, which in turn, depends on the formation temperature. The present approach of temperature estimation assumes a constant heat flux between the wellbore and formation throughout the entire operation time. However, quite often the heat transfer rate between the formation and wellbore changes with time. We used the superposition principle to account for the gradual change of heat flux with time. Analytical solutions with the assumption of invariant and linear variation of heat flux with depth, and numerical solution of the governing differential equation were obtained.

We developed expressions for fluid temperature during production, injection and mud circulation. The results showed variation in the temperature profiles when superposition is used during oil production and in mud circulation compared to solution without superposition. The solutions of linear variation of heat flux with depth assumption were close to the numercial solutions.

CHAPTER 1 INTRODUCTION

Multiphase flow is the simultaneous flow of more than one phase in a single conduit. The phases can be any combination of solids, liquids and gases. Multiphase flow is widely encountered in the petroleum industry and in the chemical process industry. It also occurs in steam generating boilers and nuclear power generators.

Each industry views multiphase flow from its own perspective. The petroleum industry has many unique features that create complications not encountered by other industries. The fluids involved are multicomponent mixtures whose phase behavior is extremely complex. The range of pressure and temperatures encountered in the petroleum industry is also very broad. It has been found that the pressure can range from 15,000 psia to near atmospheric conditions while the temperature can range from 400°F to below the freezing point of water. Pipes used in the production process from the reservoir can be either vertical or inclined. Transportation on the surface use the pipelines that are generally horizontal. Wells producing petroleum crudes can be from few hundred feet to more than 20,000 ft whereas the surface pipe can vary from a few feet to several hundred miles. Piping systems often involve significant variation in geometry, diameter, shape and inclination angle. Although most vertical and inclined systems involves cocurrent upflow,

it is not very uncommon to have downward multiphase flow in injection wells or downcomers connecting offshore platforms to subsea pipelines.

Engineers in petroleum industry are faced with the requirement to predict the relationships between flow rates and pressure drop throughout a reservoirs entire production life under different types of circumstances such as piping geometry, length, diameter, angle of inclination, etc. The pressure drops encountered during the production enter into a wide array of design calculations. The design considerations may include the tubing size and operating wellhead pressure in a flowing well; well completion or recompletion scheme; or artificial lift during gas lift or pump operation in a low energy reservoir. The pressure drop calculations are also needed in various equipment design calculation.

Simulation of multiphase flow in weils also requires the ability to predict fluid temperatures in a system that undergoes complex heat transfer between wellbore and the formation. It is essential to predict the fluid temperature with reasonable accuracy, because temperature determines various fluid properties, including the extent of dissolved hydrocarbon gases, which is a very important parameter in the process. Besides having influence on pressure profile, the temperature profile itself is also very important. For example, a very common problem in the arctic operation is due to the gas hydrate formation. The hydrates are formed at low temperature and high pressure. To design multiphase flow in a gas hydrate prone system, the phase behavior of gas hydrates, which is a function of temperature needs to be considered. Sometimes system pressure, temperature, and water contents are manipulated to avoid the gas hydrate phase envelope. Heat transfer between the wellbore and formation plays an important role under such circumstances underscoring the importance of temperature prediction.

One of the objectives of this work is to develop a unified two-phase flow model that will be uniquely useful to the petroleum industry. The other important objective is to study some of the complex heat transfer problems encountered in wellbores. We take a rigorous approach to model heat transfer in wellbores with particular attention to appropriate boundary conditions. The models will predict temperature profiles during production, injection, and mud circulation. Superposition principle is used to account for gradual change in heat transfer with time.

CHAPTER 2 THEORY

Understanding the physical behavior of multiphase flow in wells is important because hydrocarbon production, as well as well testing/production logging, often involves the simultaneous flow of two or more phases in wells that have a variety of orientation and geometry. Designing such wells tubulars requires estimation of pressure drop. For existing wells, estimating productivity or designing artificial lift also demands pressure drop calculations.

The importance of multiphase flow in chemical and petroleum industries has led to proposals of many models and correlations for pressure gradient estimation. Most of these models recognize that the in-situ gas velocity is generally higher than the in-situ liquid velocity in up flow. The higher gas velocity is caused by the buoyancy effect and the tendency of the gas phase to flow through the central portion of the channel. The difference between the two phase velocity is called slip. The in-situ gas void fraction is different than the input gas void fraction because of this slip.

The extent of the slip between the phases depend on the various configurations the phases take up depending on the prevailing conditions. These distinctive patterns make the flow pattern approach superior to entirely empirical approaches. The pioneering effort of many workers in this area has made predicting flow pattern transitions quite reliable. In this work the relationships for void fraction in terms of phase velocities and system properties are developed. In developing the models, extensive use is made of the published work in the area. As such, this work is an integration of the present knowledge on flow pattern approach in two-phase flow.

The mechanical energy balance for a flowing fluid over a differential pipe length dz without any energy input, may be written as,

$$\frac{dP}{dz} + \rho_m \frac{g}{g_c} \sin \theta + \frac{2f V_m^2 \rho_m}{g_c D} + \frac{\rho_m V_m}{g_c} \frac{dV_m}{dz} = 0$$
(2.1)

The last three terms in Equation 2.1 represent the potential energy loss, the friction loss, and the kinetic energy loss respectively. Hence, we may write the total pressure gradient, dP/dz, during single or multiphase flow as the sum of the gravitational (static head, dP/dz_H), frictional (dP/dz_F) and kinetic head (dP/dz_A) components:

$$\frac{dP}{dz} = \left[\frac{dP}{dz}\right]_{H} + \left[\frac{dP}{dz}\right]_{F} + \left[\frac{dP}{dz}\right]_{A}$$
$$= -\frac{1}{g_{c}}\left[g\rho_{m}\sin\theta + \frac{2fV_{m}^{2}}{D} + \rho_{m}V_{m}\frac{dV_{m}}{dz}\right]$$
(2.2)

The problem for two-phase flow is to find an appropriate expression for the mixture density ρ_m and the mixture friction factor f_m . For vertical flow, the static head is the major contributor to the total head loss, and in some cases. (low gas fraction and low flow rates) it may account for more than 95% of the total gradient. Since the mixture density is related to the gas void fraction E_g (in-situ volume fraction of the gas) by.

$$\rho_{m} = \rho_{g} E_{g} + \rho_{l} (1 - E_{g})$$
(2.3)

Accurate estimation of the void fraction is of paramount importance in multiphase flow analysis. The frictional head loss also requires an estimate of the mixture density and, hence, the gas void fraction. The gas void fraction depends on the in-situ velocity of the gas phase relative to the mixture. The gas phase velocity is influenced by the buoyancy effect and the tendency of the gas phase to flow through the central portion of the channel where the local mixture velocity is higher than the average velocity. Both these effects depend on the particular flow pattern that exists under the given conditions of flow, pressure and channel geometry. The various models available to estimate the void fraction and the pressure gradient are discussed in chapter 3 and 4.

Heat transfer in wellbores affects the temperature of the hydrocarbon mixture and, hence, the bubble point pressure. This in turn affect the gas volume fraction and pressure drop. Models available at present do not adequately account for heat transfer between the produced fluid and the formation. In this work, a rigorous approach has been taken to model heat transfer with particular emphasis on appropriate boundary conditions.

When a liquid is produced from a reservoir, its temperature at the bottomhole may be assumed to be same as that of the formation. While this is not true of gases, gas inlet temperature may be estimated from the formation temperature if Joule-Thompson effect is properly accounted for. Thus the bottomhole temperature of a produced fluid may be reliably estimated. However, *as* the fluid rises up the well, its temperature soon becomes significantly higher than the surrounding earth temperature because of general decline in earth temperature with decreasing depth. The temperature difference between the

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wellbore fluid and earth causes a transfer of heat from the fluid to the surrounding earth, and, therefore the fluid temperature decreases as it goes up. The transferred heat raises up the surrounding formation temperature near the wellbore. So, at any depth, the formation temperature would vary not only with the radial distance from the well, but also with production time. Hence, heat loss from the fluid decreases with time and depends on the various resistances to heat flow between the hot fluid in the tubing and the surrounding earth.

To derive an expression for fluid temperature as a function of depth and time, the formation temperature distribution needs to be established as a function of radial distance and time. An energy balance on the fluid in the wellbore can then be used to relate to the fluid temperature, the wellbore/earth interface temperature, and the heat flux between the formation and wellbore. The details of the energy balances are discussed in chapter 3 and 4.

CHAPTER 3

LITERATURE SURVEY

Design and operation of equipments involving multiphase flow often requires estimates of pressure drop within the equipment. Multiphase flow is much more complicated than single phase flow. The analysis of single phase flow is made easier if it can be established that the flow is either laminar or turbulent and whether any separation or secondary flow effect occurs. This information is equally useful in the multiphase flow, however, the geometry of the flow is of greater importance. The already intricate model developmental problem in multiphase flow is further complicated by complex heat transfer associated in the wellbores. In this chapter, we examine the various approaches presently available to estimate the void fraction and the pressure gradient in the vertical system. We also examine the heat transfer aspects associated in the wellbores.

3.1 Pressure Drop in Two-Phase Flow

Many models and correlations exist to predict pressure drop in vertical and inclined multiphase flow. Some of these were developed from large experimental data bases, relying almost entirely on empiricism. Other are mechanistically based models which are capable of accounting for the various flow patterns associated with the flow. In this section, we first discuss models based on flow pattern. Then the other major approaches are presented.

3.1.1 Flow Pattern Approach

When multiphase flow occurs, the phases take up a variety of configurations, known as flow patterns. A particular flow pattern depends on the condition of pressure, flow rate, heat flux and channel geometry. Various techniques are available for the study of two-phase flow patterns in heated and unheated channels. In transparent channels at low velocities, it is possible to distinguish the flow patterns by direct visualization. At higher velocities where the pattern becomes indistinct, flash and cine photography can be used to slow the flow down and extend the range. Numerous other ingenious techniques are also in use to examine the flow patterns. In this work only those patterns that are clearly distinguishable and generally recognized will be considered. Four such flow patterns - bubbly, slug, churn, and annular are schematically shown in Figure 1.

At low gas flow rates, the gas phase rises through the continuous liquid medium as small discrete bubbles, thus the name bubbly flow. As the gas flow rate increases, the smaller bubbles begin to coalesce forming larger bubbles. At sufficiently high gas flow rates, the agglomerated bubbles become large enough to occupy almost the entire pipe cross section, separated from the pipe wall by a thin liquid film. These large bubbles, known as Taylor bubbles, separate the liquid slugs between them. The liquid slugs, which usually contain smaller entrained gas bubbles, give the name of the flow regime. At still higher flow rates the shear stress between the Taylor bubble and the liquid film increases, which finally causes a breakdown of the liquid film and the bubbles. The





resulting churning motion of the fluids gives rise to the name of this flow pattern. The final flow pattern, annular flow, occurs at extremely high gas flow rates which causes the entire gas phase to flow through the central portion of the pipe. Some liquid is entrained in the gas core as droplets while the rest of the liquid flows up the wall through the annulus formed by the tube wall and the gas core. Models presently available for vertical system for the various flow regimes are described below.

3.1.1.A Bubbly Flow

In bubbly flow the gas phase is distributed as discrete bubbles in a continuous liquid phase. At one extreme the bubbles may be small and spherical, and at the other extreme the bubbles may be large with a spherical cap and a flat tail. In this later state, although the sizes of the bubbles do not approach the diameter of the pipe, there may be some confusion with the slug flow.

If it is assumed that during bubbly flow most of the bubbles flow through the central portion of the channel, then the in-situ velocity of the gas phase, V_g , is the sum of the terminal rise velocity, V_t , and the mixture velocity at the channel center. If the central mixture velocity is designated to be C_o times the average mixture velocity, V_m , then it can be written that,

$$V_{e} = C_{o}V_{m} + V_{i} \tag{3.1}$$

If the flow is "ideal" bubbly, which is possible at very low gas flow rates and with pure liquids, the bubbles do not affect each other's motion and Equation 3.1 is not strictly valid. In such cases, the Drift Flux model, developed by Ishii (1975), Zuber and Findlay (1965), Wallis (1969) and others, should be used. Indeed, Equation 3.1, a special form of the Drift Flux model, is valid when the bubbles are affected by the tube wall and the wakes of other bubbles. For most practical systems, fluids are rarely pure and Equation 3.1 is quite appropriate.

Noting that the in-situ velocity, V_g , of the gas phase is equal to the superficial gas velocity divided by the gas void fraction, ($V_{sg} = V_g / E_g$). Equation 3.1 may be rewritten to arrive at the following expression for the gas void fraction,

$$E_g = \frac{V_{sg}}{C_o V_m + V_i} \tag{3.2}$$

For most cases, the terminal rise velocity, V_t , appears to be well represented by the Harmathy (1960) correlation. Hasan and Kabir (1988) also suggests the use of Harmathy correlation.

$$V_{t} = 1.53 \left[\frac{g \sigma(\rho_{t} - \rho_{g})}{\rho_{t}^{2}} \right]^{0.25}$$
(3.3)

Value of the Flow Parameter C_o : Researchers analyzed various mixture velocity profiles and bubble distributions across the channel and arrived at expressions for C_o in terms of the parameters of these profiles. For most practical cases, Reynolds number based on bubble velocity is much greater than 2100. In turbulent flow the mixture velocity at the axis of the pipe is 1.2 times the average mixture velocity. If the gas bubbles are assumed to flow mostly through the central portion of the pipe, as has been shown to be the case for vertical flow, then the value of C_o is 1.2 as established in the classical work of Zuber and Findlay (1965) for an air-water system in a five cm pipe. Hasan and Kabir recommended 1.2 for the flow parameter C_o to estimate the in-situ gas void fraction during vertical bubbly flow.

Dispersed Bubbly flow : Sometimes at higher flow rates, the turbulence breaks up the larger agglomerated bubbles and the resulting flow pattern is somewhat different than the bubbly flow. This type of bubbly flow which results from the breakdown and dispersion of larger bubbles in the liquid phase is known as dispersed bubbly flow. Under certain circumstances, this is the only type of bubbly flow that can be observed in inclined system. Although slightly different, the equations developed for bubbly flow are also applicable for dispersed bubbly flow.

3.1.1.B Slug Flow

In slug flow, the gas bubbles are approximately the diameter of the pipe and are known as Taylor bubbles. The nose of the bubble has a characteristic spherical cap and the gas in the bubble is separated from the pipe wall by a slowly descending film of liquid. The liquid flow is contained in liquid slugs which separate successive gas bubbles. These slugs may or may not contain smaller entrained gas bubbles carried in the wake of the large bubble. The length of the main gas bubble can vary considerably. The pattern has also been designated by some as plug or piston flow at low flow rates where the gas liquid boundaries are well defined, and as slug flow at higher rates where the boundaries are less clear.

The analysis for slug flow is very similar to that for bubbly flow. Indeed, Equation 3.2 applies for void fraction in slug flow as well, but with different constants. Assuming there is no bubble in the liquid slug, the void fraction for ideal slug flow becomes,

$$E_{gT} = \frac{V_{sg}}{C_1 V_m + V_{tT}}$$
(3.4)

But slug flow is rarely ideal. The liquid slug contains gas bubbles in it. The cellular approach pioneered by Fernandes et al. (1983) accounts for the bubbles in the liquid slug. Hasan and Kabir simplified the Fernandes et al. (1983) approach to model slug flow. They denoted the in-situ gas fraction in the section with Taylor bubble as E_{gT} , and that in the liquid slug as E_{gs} , and obtained the following expression for average void fraction,

$$E_{g} = \left[\frac{L_{T}}{L_{C}}\right]E_{gT} + \left[\frac{L_{s}}{L_{C}}\right]E_{gs}$$
(3.5)

The terms L_s/L_c and L_T/L_c can be calculated from the following expressions,

$$E_{g} = \left[\frac{L_{\tau}}{L_{c}}\right] E_{g\tau} + 0.1 \qquad for \quad V_{sg} > 0.4 \quad m/s$$

$$E_{g} = \left[\frac{L_{\tau}}{L_{c}}\right] E_{g\tau} + 0.25 V_{sg} \qquad for \quad V_{sg} \le 0.4 \quad m/s \qquad (3.6)$$

Because the flow is almost surely turbulent, and the bubbles ride through the flat portion of the velocity profile, C_1 (as C_0 in bubbly flow) is expected to be 1.2. This is





indeed found to be the case by Nicklin et al. (1962), Hasan and Kabir (1988) and others and 1.2 is the accepted value for the parameter.

The Taylor bubble rise velocity, V_{iT}, in slug flow is given by Nicklin (1962),

$$V_{tT} = C_2 \sqrt{gD \left[\frac{\rho_t - \rho_g}{\rho_t}\right]} \simeq C_2 \sqrt{gD}$$
(3.7)

Extensive data and theoretical analyses by a number of researchers indicate that C_2 is influenced by the forces of inertia, viscosity, and surface tension. The data of White and Beardmore (1962), along with those of Dumitrescu (1943) have been represented by the following single equation by Wallis (1969),

$$C_{2} = 0.345 \left[1 - e^{-\frac{0.01 Nf}{0.345}} \right] \left[1 - e^{\frac{3.37 - Eo}{m}} \right]$$
(3.8)

where Nf is the dimensionless inverse viscosity number, $\sqrt{\{D^3g(\rho_1-\rho_g)\rho_1 / \mu_1\}}$, Eo is the Eotvos number, $gD^2(\rho_1-\rho_g)/\rho$, and m is a parameter dependent on Nf. The value of m is 10 when Nf is greater than 250, is 25 when Nf is less than 18 and is given by m = 69 (Nf)^{-0.35} for 18 < Nf < 250. For large values of Nf (say >300) and Eo (>100), Equation 3.8 reduces to C₂ = 0.345. For air-water flow through a 5 cm pipe at standard conditions, Nf = 35000 and Eo = 322. Thus, for many practical systems (if diameter is not too small) C₂ = 0.345

3.1.1.C Churn Flow

Churn flow is formed by the breakdown of the large gas bubbles in slug flow. The gas or vapor flows in more or less chaotic manner through the liquid, which is mainly displaced to the channel wall. The flow has an oscillatory or time varying character; hence the descriptive name **churn** flow. This region is also sometimes referred to as semi-annular, annular-slug transition or froth flow (Govier-Aziz (1972), Aziz-Govier-Fogarasi (1972))

The churn or froth flow pattern has not been investigated extensively because of its chaotic nature. However, the analyses presented for bubbly and slug flow should also be applicable for the churn flow pattern. Thus the equation developed for predicting void fraction in slug flow (Equation 3.7) may be used for the churn flow regime as well. Although the bubble shape is quite different from the classical Taylor bubble, the bubble rise velocity during churn flow is probably not much different from that for slug flow. In addition, because the mixture velocity is much higher than the bubble rise velocity during churn flow, a slight error in estimating V_{tr} does not significantly affect void fraction estimation. On the other hand, an accurate estimate of the flow parameter C_1 is very important for predicting void fraction. The bubble concentration profile in churn flow is unlikely to be similar to that for slug flow because of the churning motion characteristic of this flow regime. Hasan and Kabir suggested a value of 1.15 for the parameter C_1 . In this work we use Equation 3.5 and C_1 = 1.15 for the estimation of void fraction in churn flow.

3.1.1.D Annular Flow

In annular flow, the gas phase along with the entrained liquid droplets, flows through the core of the channel forming a continuous phase. The liquid phase is dragged along the pipe wall and appears to flow through the annulus formed by the channel wall and the vapor core; hence the name annular flow. Large amplitude coherent waves are usually present on the surface of the liquid film and the continuous break-up of these waves forms a source for droplet entrainment, which occurs in varying amounts in the central gas core. The droplets are separate rather than agglomerated.

In ideal annular flow, when no liquid is being carried as droplets in the gas phase and the gas-liquid interphase is smooth, the estimation of pressure drop in annular flow reduces to that of estimating pressure drop in single phase gas flow. The liquid film thickness is typically less than 5 % of the tube diameter, thus introducing little error even if it is neglected in calculating the channel diameter for gas flow.

Unfortunately, however, annular flow is rarely ideal. Usually, a substantial fraction of the liquid is carried as droplets in the gas stream requiring estimation of the mixture density. In addition, the gas-liquid interface is usually wavy and determining the appropriate friction factor becomes very difficult.

The following equation may be used for the total pressure gradient during annular flow noting that V_g replaces V_{sg} .

$$-\frac{dP}{dz} = \frac{1}{g_c} \left[\frac{2f_f V_g^2 \rho_c}{D} + g\rho_c + \rho_c V_g \frac{dV_g}{dz} \right]$$
(3.9)

The acceleration term in this equation contains the differential dV_g/dz . This term can be rewritten in terms of dP/dz and V_g using the gas law and thereby Equation 3.9 becomes

The problem then reduces to that of estimating the density of the fluid in the core, ρ_c , and estimating the friction factor f_f . To determine the density of the fluid flowing

$$-\frac{dP}{dz} = \frac{1}{g_c} \left[\frac{\frac{2 f_f V_s^2 \rho_c}{D} + g \rho_c}{1 - \frac{\rho_c V_s^2}{P g_c}} \right]$$
(3.10)

through the core it is necessary to estimate the entrainment. Hasan and Kabir (1988) recommended the following correlations proposed by Wallis and Steen (1964) for the estimation of entrainment.

$$E = 0.0055 \left[10^4 (V_{sg})_c \right]^{2.86} \qquad if \quad 10^4 (V_{sg}) < 4$$
$$E = 0.857 \log_{10} \left[10^4 (V_{sg})_c \right] - 0.20 \qquad if \quad 10^4 (V_{sg})_c > 4 \qquad (3.11)$$

A number of correlations are available for predicting the film friction factor f_f . Hasan and Kabir (1988) recommended the one proposed by Wallis (1969), which probably is the best among these.

$$f_{f} = f_{g} \left[1 + 75 \left(1 - E_{g} \right) \right]$$
(3.12)

Some rigorous models, which incorporate velocity profile in the liquid and gas core have been developed in recent years. Considering the rarity of this flow regime in oilwell and the complexity of those models, we use the simple approach presented here.

3.1.1.E Transition criteria

The individual models discussed so far enables us to estimate void fraction and the pressure gradient once the flow pattern is established. But it is very difficult to correctly determine exactly when transition from one flow pattern to another takes place. One reason behind this problem is the lack of agreement in the description of the flow pattern. Besides, the transition does not occur abruptly. In most cases gradual transition from one pattern to another is observed.

Bubbly-Slug Transition. Transition from the condition of small bubbles dispersed throughout the flow cross-section to the condition when the bubbles become large enough to fill almost the entire pipe cross-section, requires a process of agglomeration or coalescence. In general, bubbles, other than very small ones, follow a zig-zag path when rising through the liquid. This results in collision between the bubbles, with consequent bubble agglomeration and formation of larger bubbles. Obviously, collision frequency and bubble agglomeration increases with increasing gas flow rates. Radovich and Moissis (1962) theoretically examined the behavior of bubbles by considering a cubic lattice in which the individual bubbles fluctuate. It was found that at a void fraction of 0.3, the collision frequency becomes so high that a transition to slug flow is to be expected. Griffith and Snyder (1964) experimentally verified that the transition occurs at a void fraction of 0.25 to 0.3. Hasan and Kabir also found the transition to take place at a void fraction of about 0.25.

Thus $E_g = 0.25$ may be taken as the criteria for transition between bubbly and slug flow. This criteria when expressed in terms of the superficial velocities by equating the slip between the phases with the terminal rise of a single bubble then the relationship between V_{sg} and V_{sl} at the transition becomes.

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$$V_{sg} = 0.429 V_{sl} + 0.357 V_{l}$$
(3.13)

Since the transition from bubbly to slug flow is likely to be gradual, it is unlikely, although assumed in deriving the above equation that the void fraction relationship for bubbly flow would be applicable up to the point of transition. The appropriate expression in slug flow is similar to that of the bubbly flow with only exception in bubble rise velocity. However, the difference between V_{tT} and V_t is not large and the above equation is adequate in representing the transition between bubbly and slug flows.

Dispersed Bubbly Flow : The transition criteria discussed above applies only at low or moderate flow rates. At higher flow rates, the turbulence breaks up the larger agglomerated bubbles and inhibits transition to slug flow. The bubbly flow may persist even when the void fraction exceeds 0.25 in this case. Taitel et al. (1980) analyzed the onset of dispersed bubbly flow based on the maximum bubble diameter possible under highly turbulent conditions. They concluded that if the turbulence is high enough so that the bubbles are smaller than the critical diameter, agglomeration is suppressed and bubbly flow continues. They derived the following minimum mixture velocity for dispersed bubbly flow.

$$V_{m} = 4 (D)^{0.43} \left[\frac{(\rho_{l} - \rho_{g}) g}{\sigma} \right]^{0.446} \left[\frac{\rho_{l}}{\mu_{l}} \right]^{0.072} \left[\frac{\sigma}{\rho_{l}} \right]^{0.446}$$
(3.14)

If the mixture velocity is higher than that given by the above equation, bubbly flow will persist even if the void fraction is higher than 0.25. However, it was found that bubbly flow can not persist above a void fraction of 0.52. At higher void fractions, transition to either slug or churn flow will occur.

Slug-Churn Transition. A characteristic of slug flow is the liquid confined between the Taylor bubble and the tube wall. This liquid flows around the bubble as a falling film. The interaction between this falling film and the Taylor bubble increases with increasing flow rate. The upper limit of slug flow occurs when the interaction becomes high enough to break up the bubbles, causing transition to churn flow.

The most promising model for this transition appears to be the one proposed by Brauner and Barnea (1986). They analyzed the condition of the liquid slug following the Taylor bubble just before the transition to churn flow takes place. The normal upper limit of local gas void fraction in the liquid slug is about 25%, because at higher gas fraction, the smaller bubbles coalesce to give rise to more Taylor bubbles. If the mixture velocity is high enough for dispersed bubbly flow, the local in-situ gas volume fraction in the liquid slug could attain a maximum value of 52%. Thus Brauner and Barnea argue that the transition to churn flow occurs when the void fraction in the liquid slug, which would be approximately the same as the average void fraction in the pipe, is over 52% and the mixture velocity is high enough to sustain dispersed bubbly flow. They assumed that at high flow rates void fraction may be approximated by input volume fraction so that $E_g=V_{sg}/V_M$. Hence at transition from slug to churn flow,

$$\frac{V_{s_{g}}}{V_{s_{g}} + V_{sl}} > 0.52 ; \quad Hence, \quad V_{s_{g}} > 1.08 \ V_{sl}$$
(3.15)

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Transition to Annular Flow. At high gas flow rates, transition from churn and slug flow to annular flow takes place. The liquid flows upward along the tube wall, while the gas flows through the center of the tube. The liquid film has a wavy interface and the waves could break away and be carried away as entrained droplets.

One approach to study the transition from churn (or slug) to annular flow is by analyzing the minimum gas flow rate required to reverse the direction of flow of a falling liquid film (Wallis (1969), Jones and Zuber (1978)). Another approach, adapted by Taitel et al. (1980), is to examine the drag force necessary to keep the entrained liquid droplets in suspension during annular flow. When the gas velocity is not sufficient to keep the liquid droplets in suspension, the droplets will fall back, accumulate, form a bridge, and finally establish churn or slug flow. The minimum velocity required to keep the droplets in suspension may be determined from a balance of the drag forces on these droplets and the gravitational forces acting on them

$$V_g = \frac{2}{\sqrt{3}} \sqrt{\frac{g(\rho_l - \rho_g)d}{\rho_g C_d}}$$
(3.16)

Substituting the droplet diameter d in the above equation by the maximum stable drop size, Taitel et al. (1980) arrived at the following minimum gas velocity for transition to annular flow:

$$V_{s_{\mathcal{R}}} = 3.1 \left[\frac{g \sigma(\rho_{l} - \rho_{\mathcal{R}})}{\rho_{\mathcal{R}}^{2}} \right]^{0.25}$$
(3.17)

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3.1.2 Other Approaches

A number of procedures have been reported in the petroleum engineering literature that attempts to predict pressure drop in vertical wells over a broad range of multiphase flow conditions. Four overall predictive schemes are described below because these are well known to the petroleum engineers. They are due to (1) Orkiszewski (1967), (2) Aziz, Govier and Fogarasi (1972), (3) Duns and Ros (1963), and (4) Beggs and Brill (1973).

3.1.2.A Orkiszewski Method

In 1967, Orkiszewski examined the available correlations for predicting multiphase pressure drop in vertical wells in light of data from 148 wells. He proposed a composite method based on the flow pattern approach. He recognized four different flow patterns - bubbly, slug, transition (churn) and mist (annular). To estimate void fraction and pressure drop in bubbly flow, he accepted the suggestion of Griffith and Wallis (1961). The bubbly to slug flow transition is also given by Griffith and Wallis (1961), while the transition from slug to churn and churn to annular is given by the criteria suggested by Duns and Ros (1963).

The Orkiszewski method was an improvement over the methods generally used in the petroleum industries. At present, however, simpler models with better theoretical basis and greater accuracy are available.
3.1.2.B Aziz, Govier and Fogarasi Method

In 1972, Aziz et al. presented a procedure for predicting pressure drop in vertical oil wells by combining the available literature in the area. The method used the flow pattern map of Govier et al (1957). It restricts itself in developing a prediction procedure for bubbly and slug flow only, perhaps because of the rarity of the other two patterns in oil wells.

The Aziz et al. model is somewhat similar to the Hasan Kabir model. The difference between the prediction and actual data in the bubbly flow regime is slight, but the difference is large in slug flow, when the taylor bubble rise velocity is different from the terminal rise velocity of small bubbles. Moreover, this method neglects the acceleration term in the estimation of total pressure gradient.

3.1.2.C Duns and Ros Method

In the early sixties, Ros (1961) and Duns and Ros (1963) developed a general empirical correlation from a large set of laboratory data. The method is flow regime based, but the regime definitions are somewhat different from present standard definitions. They define region I as the flow regime where the liquid is the continuous phase, and hence, include bubble, froth (presumably dispersed bubbly), plug, and some slug flow. Region II covers situations when neither phase is continuous, and hence, include the rest of the slug flow and froth flow as well as 'heading' (or pulsating flow). When gas becomes the continuous phase, as in annular-mist flow, it is termed region III. Duns and Ros also include a transition region (probably corresponding to churn flow) between Region II and Region III. The prediction by this method is quite accurate. However, its drawback is its non-standard flow pattern descriptions and transition criteria. It also contains a large number of empirically determined constants and its entirely empirical nature makes interpolation and extrapolation risky.

3.1.2.D Beggs and Brill Method

The classical study of Beggs and Brill probably gives the most comprehensive method available for predicting void fraction and pressure drop in inclined systems. Their correlation is based on a predictive method for the horizontal system and modifications to account for the inclination of the system. For estimating liquid holdup for a horizontal system, $E_{1,90}$ (= in situ liquid fraction = 1 - $E_{g,90}$), they propose the following equation in terms of mixture Froude number, Fr_m (= V_m^2/gD) and the input liquid volume fraction E_{li} .

$$E_{l,90} = a \frac{E_{li}^{b}}{(Fr_{\pi})^{c}}$$
(3.18)

The values of the parameters a, b and c depend on the flow regime. For inclined systems, Beggs and Brill use the holdup calculated by Equation 3.18 and multiplies it by an inclination factor, $F(\theta)$. The value of the multiplier depends upon the pipe inclination, input liquid fraction, dimensionless liquid velocity number, the Froude number and the flow pattern that would exist in a equivalent horizontal system.

The predictions of the Beggs and Brill correlation are usually good for inclined systems. However, the complications involved in the calculation procedure and the methods exclusive reliance on empiricism, makes it less than completely satisfactory. One problem with the correlation is that liquid input fraction, E_{li} , is used to determine the horizontal flow pattern and the correction factor, $F(\theta)$. For stagnant liquid columns, when E_{li} is zero, the method cannot be used, and for small values of E_{li} , the predictions of the method would be unreliable.

3.2 Heat Transfer in Wellbores

The importance of various aspects of heat transfer between a wellbore fluid and the earth has generated a rich literature on the subject. The usefulness of fluid temperature measurement was pointed out as early as 1937 by Schlumberger et al. (1937). Fluid temperature depends on heat loss from the wellbore to the surrounding formation. The formation temperature distribution around a well was first modeled by Ramey (1962). He neglected the effect of kinetic energy and friction, and the model was applicable only to the flow of single phase fluids. Moreover, his assumption of a vanishingly small well radius in solving the formation temperature distribution proved untenable in some cases. Many other researchers have suggested various procedures for estimating wellbore fluid temperature. The following section discusses one such model, with appropriate initial and boundary conditions.

3.2.1 Formation Temperature Distribution

Heat diffusion in a three-dimensional solid may be mathematically treated as a two-dimensional problem if symmetry around the heat source (or sink) is assumed, as in the case of a producing or an injection well. In addition, if small increment of the vertical direction of the well is considered, the problem simplifies to one-dimensional heat diffusion, because vertical heat diffusion can be ignored due to small vertical temperature gradient. This approach proposed by Hasan and Kabir (1991) has been adapted in this work. It introduces very little error and allows analytical solution of the problem which is often preferable to the alternative of tedious and time consuming numerical solution. A number of interesting heat conduction problems of similar nature was earlier presented by Carslaw and Jaeger (1950).

In a short time-step, the heat flux from the wellbore may be assumed to remain constant at a given depth. An energy balance on the formation then leads to the following partial differential equation derived in cylindrical coordinates for the variation of formation temperature with radial distance from the well and time of production:

$$\frac{\partial^2 T_{\epsilon}}{\partial r^2} + \frac{1}{r} \frac{\partial T_{\epsilon}}{\partial r} = \frac{c_{\epsilon} \rho_{\epsilon}}{k_{\epsilon}} \frac{\partial T_{\epsilon}}{\partial t}$$
(3.19)

where T_e is the temperature of earth at time t and radial distance r measured from the center of the wellbore. c_e , ρ_e and k_e are the heat capacity, density and thermal conductivity of formation. This equation is analogous to the pressure diffusion equation as used in the pressure transient literature.

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Initially, formation temperature at any given depth is constant, leading to the following condition,

$$\lim_{\epsilon \to 0} T_{\epsilon} = T_{\epsilon} \tag{3.20}$$

At the infinite or outer boundary, formation temperature does not change with radial distance, i.e.,

$$\lim_{r \to \infty} \frac{\partial T_r}{\partial r} = 0 \tag{3.21}$$

The other boundary condition is derived from the heat flow rate at the interface of wellbore and the formation, which is governed by Fourier's law of heat conduction. Rate of flow of heat per unit mass of wellbore fluid per unit length of the well, ϕ , is then given by,

$$\phi = -\frac{2\pi k_{e}}{W} \frac{r \partial T_{e}}{\partial r} \Big|_{r \text{ at wellbore}}$$
(3.22)

where W is the wellbore fluid mass flow rate, and r_{wb} is the outer radius of the wellbore.

To facilitate solution and to have more general applicability of the solution, the above equations were first recast in dimensionless variables and the solution was carried out using Laplace transformation (Lok (1991), Hasan and Kabir (1991)). The analysis resulted in the following expression for formation temperature as a function of radial distance and time.

The above equation shows that the computation of formation temperature requires evaluation of an integral involving modified Bessel function of zero and first order over

where,
$$I = \int_{0}^{\infty} \frac{1 - e^{-u^{2} t_{D}} Y_{1}(u) J_{0}(ur_{D}) - J_{1}(u) Y_{0}(ur_{D})}{u^{2} - J_{1}^{2}(u) + Y_{1}^{2}(u)} du$$
 (3.23)

the limits of zero to infinity for the dummy variable, u. Such computations are time consuming. Hasan and Kabir found that the following expressions approximate the actual solution quite reasonably.

$$T_{D} = 1.1281 \sqrt{t_{D}} \left[1 - 0.3 \sqrt{t_{D}} \right] \qquad if \ t_{D} \le 1.5$$

$$T_{D} = \left[0.4063 + 0.5 \ln(t_{D}) \right] \left[1 + \frac{0.6}{t_{D}} \right] \qquad if \ t_{D} > 1.5 \qquad (3.24)$$

where,

$$t_{D} \equiv \frac{\alpha t}{r_{wb}^{2}} ; \qquad T_{D} \equiv -\frac{2\pi k_{e}}{W\phi} (T_{wb} - T_{ei}) \qquad (3.25)$$

3.2.2 Wellbore Fluid Energy Balance

Heat loss experienced by the fluid as it flows up the well results in lowering of its temperature. An energy balance on the fluid may be done following any standard text on thermodynamics. Ramey (1962) made an energy balance on the fluid by assuming single-phase flow. Hasan and Kabir developed a more rigorous balance for a differential lengt¹, dz, for a two-phase system,

$$\frac{dH}{dz} + \frac{g\sin\theta}{g_c J} + \frac{V}{g_c J} \frac{dV}{dz} = \frac{dq}{dz} = \phi \qquad (3.26)$$

where ge and J represent appropriate conversion factors.

Fluid enthalpy, H, depends on its pressure and its temperature, which allows to write the following expression for dH/dz,

$$\frac{dH}{dz} = \left(\frac{\partial H}{\partial P}\right)_{T_f} \frac{dP}{dz} + \left(\frac{\partial H}{\partial T_f}\right)_p \frac{dT_f}{dz}$$
$$= -C_f c_p \frac{dP}{dz} + c_p \frac{dT_f}{dz}$$
(3.27)

where C_J is the Joule-Thompson coefficient and c_p is the heat capacity of the fluid at constant pressure. Combining Equations 3.26 and 3.27,

$$\frac{dT_f}{dz} = \frac{1}{c_p} \frac{dH}{dz} + C_J \frac{dP}{dz}$$
$$= C_J \frac{dP}{dz} + \frac{1}{c_p} \left[\phi - \frac{g \sin \theta}{g_c J} - \frac{V}{g_c J} \frac{dV}{dz} \right]$$
(3.28)

The radial heat transfer between the fluid and the surrounding earth may be expressed in terms of an overall heat transfer coefficient following any standard text on heat transfer (McAdams, 1942) or on transport phenomena (Bird, Stewart, and Lightfoot, 1960). Ramey (1962) and Willhite (1967) presented detailed discussions which lead to the following equation for heat transfer rate, ϕ , and the wellbore temperature, T_{wb} ,

$$\phi = -\frac{2\pi r_{io} U_{io}}{W} (T_f - T_{wb})$$
(3.29)

The overall heat transfer coefficient based on tubing outside surface area, U_{ω} , depends on resistance to heat flow from the tubing fluid to the surrounding earth and is discussed in detail in the appendix. In general, resistances to heat flow through the tubing or casing metal may be neglected. Usually natural convection is considered as the dominant heat transfer mechanism for the fluid in annulus. Resistance through the cement layer can be important depending on its thickness.

Using the definition of dimensionless temperature, T_D , in Equation 3.21, an expression for heat transfer from the wellbore/earth interface to the earth can be obtained,

$$\phi = -\frac{2\pi k_{\epsilon}}{WT_{p}} \left(T_{wb} - T_{\epsilon i} \right)$$
(3.30)

Combining Equations 3.29 and 3.30 to eliminate the wellbore temperature, T_{wb} ,

$$\phi = -\frac{2\pi}{W} \left[\frac{r_{io} U_{io} k_{\epsilon}}{k_{\epsilon} + T_D r_{io} U_{io}} \right] \left(T_f - T_{\epsilon i} \right)$$
(3.31)

3.2.3 Wellbore Fluid Temperature Distribution

Hasan and Kabir (1991) obtained an expression for variation of fluid temperature with well depth by substituting the expression for ϕ from Equation 3.31 into Equation 3.28.

$$\frac{dT_f}{dz} = \frac{T_{ei} - T_f}{A} - \frac{g}{g_c} \frac{\sin\theta}{Jc_p} + C_f \frac{dP}{dz} - \frac{V dV}{g_c J c_p}$$
(3.32)

where,
$$A = \frac{c_p W}{2 \pi} \left[\frac{k_e + (r_{io} U_{io} T_D)}{r_{io} U_{io} k_e} \right]$$
(3.33)

The undisturbed earth temperature, T_{ei} , is generally assumed to vary linearly with depth. Thus,

$$T_{e} = T_{eibh} - g_T z \tag{3.34}$$

where g_T represents the geothermal gradient and T_{eibh} is the undisturbed (static) earth temperature at the bottomhole. Even when different geologic formations are encountered at various depths, the computation may be divided into a number of zones with constant geothermal gradient being applied to each zone. If we assume that the sum of the last two terms in Equation 3.32 does not vary with well depth then Equation 3.32 becomes a linear differential equation,

$$\frac{dT_f}{dz} = \frac{T_{ei} - T_f}{A} - \frac{g}{g_c} \frac{\sin\theta}{Jc_p} + \sigma$$
where $\sigma = C_J \frac{dP}{dz} - \frac{V dV}{g_c J c_p}$
(3.35)

Equation 3.35 may be integrated for a constant A and appropriate boundary conditions. Thus, for a producing well at the bottomhole condition $(z = z_{bh})$, fluid and earth temperatures are generally known ($T_f = T_{fbh}$, and $T_{ei} = T_{eibh}$), giving the following expression for fluid temperature as a function of well depth and production time,

$$T_{f} = T_{e} + A \left(-\frac{g}{g_{e}} \frac{\sin\theta}{Jc_{p}} + \sigma + g_{T} \right) .$$

+ $e^{(c_{m}-z)/A} \left[T_{fwh} - T_{ewh} + A \left(\frac{g}{g_{e}} \frac{\sin\theta}{Jc_{p}} - \sigma - g_{T} \right) \right]$ (3.36)

The value of the parameter, σ , used in Equation 3.36 would depend on a number of variables such as flow rates, gas/liquid ratio. wellhead pressure, etc. In their work, Hasan and Kabir used the empirical expression for σ developed by Sagar et al. (1989). It should be pointed out that Equation 3.35 may be integrated for other conditions also. For example, for an injection well the wellhead fluid and earth temperatures are used as boundary conditions. In addition, for gas-lift with gas injection at known depths, or formations with numerous zones with different properties, such as geothermal gradients or conductivities. Equation 3.35 may also be integrated separately for each section, using the fluid temperature calculated at the previous section as the input for the next one.

CHAPTER 4

PROPOSED APPROACH

One of the objective of this work is to develop a multiphase flow simulator which can predict pressure profile under different type of circumstances. This thesis uses the basic model of Hasan and Kabir which has been described in chapter 3. In the first section of this chapter, we describe modifications that are needed to use the basic model for other systems. The second and third sections deal with the heat transfer aspects in the wellbores. The concept of varying heat flux is introduced first and then its application to different systems are discussed.

4.1 Unification of Two-Phase Flow Model

We have already discussed the Hasan-Kabir model to estimate the void fraction and pressure drop when the flow is in the vertical upward direction. In the following three sections, we show how the same model with some modifications can be used for the inclined systems, liquid-liquid systems, countercurrent systems, systems where flow is in downward direction and also to the conduits other than conventional circular ones.

4.1.1 Inclined Flow

The proposed unified model utilizes the flow pattern approach of Hasan and Kabir with modification for the system deviation from vertical orientation. It should be noted that for annular and dispersed bubbly flow the flow rates are very high. Consequently, the influence of buoyancy is small, and hence the effect of pipe inclination is negligible. Therefore, for the annular and the dispersed bubbly flow regimes, the relationships developed for vertical system can be used without any modification. However, this two flow patterns are not very common in petroleum production. This leaves the bubbly and intermittent flow patterns, for which the predictive scheme is described below.

Bubbly Flow. The procedure for estimating void fraction, and pressure drop in inclined system, is similar to the vertical flow. However, we would need values of flow parameter C_o and bubble rise velocity V_t for an inclined system. For vertical systems, we are able to reason that the value of flow parameter C_o should be 1.2 because the flow is turbulent and the bubbles ride the central portion of the channel where the mixture velocity is 1.2 times the cross-sectional average velocity. One would expect the value of the parameter C_o to be influenced by the deviation of the pipe, since the bubble concentration profile would be affected by the pipes inclination. But, this effect has been experimentally observed to be very small, and the value of C_o has been generally found to be 1.2. In addition, the bubble rise velocity has been found to remain unchanged with pipe inclination.

In a vertical system, the transition from bubbly to slug flow occurs at a void fraction of about 0.25. This criteria for the transition from bubbly to slug flow should also be applicable to inclined systems. However, in an inclined pipe the gas phase tends to flow along the upper wall of the pipe, thereby increasing the actual local void fraction. Conceivably, this local void fraction in the upper section of the channel may exceed 0.25, even when the cross-sectional average void fraction is much smaller than 0.25. As a result, in an inclined pipe transition to slug flow occurs at a cross- sectional average void fraction of less than 0.25.

For vertical systems, void fraction in bubbly flow is given by,

$$E_{g} = \frac{V_{sg}}{C_{o}(V_{sl} + V_{sg}) + V_{t}}$$
(4.1)

This relationship may be applied locally in the case of an inclined pipe if the actual superficial velocity of the gas phase at the upper section of the pipe is used rather than the cross-sectional average value. In an inclined pipe, it is reasonable to assume that the actual cross-sectional area available for the gas to flow is the projection of the cross-sectional area on a horizontal plane. The local superficial velocity of the gas phase therefore, is $(V_{sg})_{loc} = q_g/A \sin \theta = V_{sg}/\sin \theta$. Using $(V_{sg})_{loc}$ in place of V_{sg} in the void fraction relationship, we obtain the local void fraction at the upper portion of the pipe wall, $E_{g, loc}$,

$$E_{g,loc} = \frac{\left[V_{sg} / \sin \theta \right]}{C_o \left[V_{sl} + (V_{sg} / \sin \theta) \right] + V_t}$$
(4.2)

Rearranging and using $E_g = 0.25$ and $C_o = 1.2$ at transition,

$$V_{st} = \left[0.43 \, V_{st} + 0.357 \, V_{t} \right] \sin \theta \tag{4.3}$$

Slug Flow. Hasan and Kabir (198?) found that their model for vertical upward flow can be used for inclined systems with some modifications. The models remain almost the same, but value of some of the parameter changes. The value of C_1 in vertical flow, was taken as 1.2. Experimental verification of this value for C_1 for inclined systems has been provided by Patel (1986). However, the terminal rise velocity of a Taylor bubble is significantly influenced by the pipe inclination. This fact is evident from the classical work of Runge and Wallis (1965) and Zukoski (1966). Their data generally indicate that the Taylor bubble rise velocity increases with increasing deviation of the pipe from vertical, until a maximum is reached for a deviation angle of about 50°. The terminal rise velocity then gradually decreases with increasing deviations and finally becomes zero for horizontal systems.

An expression for the rise velocity of a bubble may be derived by balancing the buoyancy force against the drag force experienced by a rising bubble. Such an expression for the rise velocity of a Taylor bubble in an inclined pipe, $V_{cT\theta}$, can be derived in the in terms of the rise velocity in a vertical pipe, V_{cT} , and the angle of inclination θ .

$$V_{dT\theta} = V_{dT} \sqrt{(\sin\theta)(1 + \cos\theta)^{1.2}}$$
(4.4)

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4.1.2 Downward Flow

Downward simultaneous flow of gas and liquid, although less common than twophase up-flow, is important in chemical process industries and petroleum crude production. Wet steam injection into high viscosity oil bearing formations is an example of the two phase down-flow. As in the up-flow, the static head is quite often the major contributor to the total head loss, especially for vertical and near vertical systems. Consequently, an accurate estimation of gas void fraction, E_g , is required because the mixture density is related to the void fraction.

For up-flow, the effect of buoyancy and the tendency of the gas phase to flow through the channel center causes in-situ gas velocity to be higher than the mixture velocity. For down-flow, buoyancy will oppose the downward flow of the gas phase. The cross-sectional distribution of the gas phase in the channel may also be different from that in the up-flow. The effect of buoyancy and bubble distribution across the flow channel also depend on the existing flow pattern. The relationship for void fraction in terms of phase velocities and system properties and the upper limit for the flow regime is described in this section.

Bubbly Flow. Equation 4.1 would also apply to downward bubbly flow in vertical and inclined systems. But, because in downward flow the terminal rise velocity acts opposite to the direction of flow, the expression for void fraction can be written as,

$$E_{g} = \frac{V_{sg}}{C_{o}(V_{sl} + V_{sg}) - V_{l}}$$
(4.5)

Hasan (1989) found that the Harmathy correlation applicable to the bubble rise velocity in both vertical and inclined pipes. He also found that a value of 1.2 for the flow parameter C_o to agree with the data of Mukherjee (1978) and Beggs (1972). The same value has been used in this work.

The transition from bubbly to slug flow has been found to occur at a void fraction of about 0.25 in upward flow. This criteria would apply to downward flow as well. Using a void fraction of 0.25 Hasan and Kabir arrived at the following expression,

$$V_{sg} = \left[0.43 \, V_{sl} - 0.36 \, V_{l} \right] \sin \theta \tag{4.6}$$

Slug Flow. Slug flow is characterized by a Taylor bubble that fills up almost the entire pipe cross-section followed by a liquid slug that contains small gas bubbles. Hasan (1989) simplified the cellular approach of Hasan and Kabir (1988) for vertical up-flow for adaptation to down-flow. The analysis is similar to that of up-flow, except that the terminal rise velocity acts opposite to the mixture flow direction.

Taylor Bubble Rise Velocity : In upward flow the variation in Taylor bubble rise velocity with pipe inclination has been given by Hasan and Kabir. The same expression can be successfully used in downward flow.

Flow Parameter C_o : Hasan (1989) tried different values for the parameter C_o to fit the experimental data of Mukherjee (1978). It was been found that $C_o=1.12$ serves the purpose quite reasonably. Therefore, a constant value of 1.12 for C_o has been used for the downward slug flow in this model.

4.1.3 Flow in Different Geometries

The majority of two-phase flow occurs in circular conduit. But flow through other geometries, especially through annuli is very common. This section deals with the flow pattern transition and void fraction estimation in non-circular channels.

Flow Through Annuli. The presence of an inner tube does not appear to influence the bubble concentration profile in bubbly flow. The value of the flow parameter C_o , for annuli has been found to remain essentially the same as that for the circular channel. Bubbly-slug transition was found to take place at the same void fraction of 0.25 in the annular geometry. Thus, the transition criteria remains the same as that of circular channels. The dispersed bubbly model described for the circular channels applies to the annuli also.

Although the presence of a inner tube does not affect the bubbly or dispersed bubbly flow, but a significant effect is found in slug flow. The nose of the Taylor bubble becomes sharper which causes an increase in the rise velocity V_{tT} . We use the following expression suggested by Hasan and Kabir for estimation of Taylor bubble rise velocity.

$$V_{tTa} = \left[0.345 + 0.10 (D_i/D_o) \right] \left[\sqrt{g D_o (\rho_i - \rho_g)/\rho_i} \right]$$
(4.7)

where D_i and D_o represents the tube and annular diameter.

Sadatomi et al. (1982) defined 'Equi-peripheral' diameter as the wetted perimeter of the channel divided by π , which is $D_i + D_o$ for the annulus and used that in estimation of Taylor bubble rise velocity. The equi-peripheral diameter is different than the equivalent diameter which is $D_o - D_i$. But the agreement of Sadatomi et al. correlation with

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their data from annuli is less satisfactory than data with other channels. We use Equation 4.7 in our model.

The slug-churn and churn-annular transition remains similar to that of circular channels. Void fraction and pressure gradient can be estimated in a fashion similar to that for circular channels.

Flow Through Other Geometries. Sadatomi et al. (1982) found that the geometry of the pipe does not influence the bubble concentration profile considerably. The models for circular conduits can be used with other geometries also. The diameter of the circular pipe should be replaced by the equi-periphery diameter which is expressed in terms of periphery, $D_e = Periphery / \pi$.

4.1.4 Countercurrent Flow

Countercurrent two-phase flow is encountered occasionally in oil and gas production, in well testing, and in production logging. Little research have been done to understand countercurrent system, where the liquid flows downward while the gas moves upward.

The models for two-phase flow described in chapter 3 can be used for countercurrent system with some modifications. The mixture velocity is the difference between the gas and liquid superficial velocity instead of their sum used in earlier situations. Recognizing this modification, the void fraction in **bubbly flow** becomes,

$$E_{g} = \frac{V_{sg}}{C_{o} (V_{sg} - V_{sl}) + V_{t}}$$
(4.8)

A value of 2.0 for C_o and Harmathy equation for V_t has been found to well represent the experimental data (Srinevasan (1993)).

Transition from bubbly to slug is also expected at a void fraction of 0.25 during the countercurrent flow. Using this value for void fraction the following transition criteria in terms of superficial velocity is obtained,

$$V_{sg} = \frac{V_t - C_o V_{sl}}{4 - C_o} \sin \theta$$
(4.9)

The void fraction in slug flow can be calculated using the general approach described in chapter 3 with the modifications noted for the bubbly flow region. The terminal rise velocity V_t should be replaced by the Taylor bubble rise velocity, V_{tT} , and C_o value changes from 2.0 to 1.2. The churn and annular flow regions are rarely observed in countercurrent situations and little has been understood about their behavior.

4.1.5 Liquid - Liquid Flow

Liquid-liquid two-phase flow is commonly encountered in chemical process industries and is quite prevalent in the production of petroleum crudes. Although a lot of work has been done to understand the gas-liquid two-phase flow, few investigators have attempted to explain the mechanics of simultaneous flow of two immiscible liquids. The physics of liquid-liquid flow is different from that of the gas-liquid flow. However,

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there are some similarities between them as well. The model for two phase flow described earlier can be used for liquid-liquid system with some modifications.

The in-situ volume fraction of the lighter phase (i.e. oil in a mixture of oil-water) depends on its in-situ velocity relative to the mixture. The in-situ oil velocity, V_o , is influenced by the tendency of the oil droplets to flow through the central portion of the channel where the local mixture velocity is greater than the cross-sectional average velocity. We use the approach suggested by Hasan and Kabir (1987) and Wallis (1969). According to them, the density difference between phases give rise to drift flux, j_{ow} , which adds velocity to the lighter phase. Hence, lighter phase velocity becomes,

$$V_o = C_o V_m + \frac{j_{ow}}{E_o}$$
(4.10)

where $j_{ow} = V_t E_o (1-E_o)^2$, as suggested by Wallis (1969).

For ideal bubbly flow, when the bubbles do not interact, taking Wallis suggestion into account, the following expression for lighter fluid volume-fraction is obtained,

$$E_o = \frac{V_{so}}{1.2 V_m + V_t (1 - E_o)^2}$$
(4.11)

Hasan and Kabir (1987) found Harmathy equation suitable in estimating the terminal rise velocity in liquid-liquid systems. The following expression represents the transition from bubbly to slug flow,

$$V_{so} = 0.43 V_{sw} + 0.20 V_{t}$$
(4.12)

The expressions derived for bubbly flow are also applicable for slug and churn flow. However, Hasan and Kabir suggested to change the value of the flow parameter C_o from 1.2 to 1.15 for churn flow. Mist flow, analogous to annular flow in gas-liquid system is less frequent and should be treated as homogeneous flow.

4.2 Heat Transfer During Production and Injection

The steps involved in formulating an expression for fluid temperature has been described in detail in chapter 3. One of the underlying assumptions in the process was that the heat flux from the formation to the fluid remained constant throughout the entire production time. In order to estimate the fluid temperature more accurately it is essential to incorporate changes that will account for the variation of heat flux with production time. We will develop the concept of varying heat flux in the following section and use it in subsequent sections.

4.2.1 Effect of Varying Heat Flux

The rate of heat transfer, ϕ , from the wellbore to the formation (or vice versa) at the formation/wellbore interface per unit depth of the well is given by

$$W \phi = -\frac{2\pi k_e}{T_p(t_p)} \left(T_{wb} - T_{ei} \right)$$

$$(4.13)$$

The dimensionless temperature, $T_D(t_D)$, is a function of dimensionless time, $t_D = \alpha t/r^2$, and can be easily estimated from (Hasan and Kabir, 1991),

$$T_{D} = 1.1281 \sqrt{t_{D}} \left[1 - 0.3 \sqrt{t_{D}} \right] \qquad if \quad t_{D} \le 1.5$$

$$T_{D} = \left[0.4063 + 0.5 \ln(t_{D}) \right] \left[1 + \frac{0.6}{t_{D}} \right] \qquad if \quad t_{D} > 1.5 \qquad (4.14)$$

However, Equation 4.13 is only valid for constant heat flux at the wellbore/formation interface. In general, fluid temperature within the wellbore tends to approach the temperature of the formation surrounding it, thereby decreasing heat transfer rate with time. It is possible to account for this changing heat flux by using the superposition principle. Lets consider a new well that has produced fluids at a constant rate for a time t. To estimate fluid temperature at time t, we divide the total into n periods (not necessarily equal) – $(t_1 - 0)$, $(t_2 - t_1)$, $(t_3 - t_2)$, ..., $(t_{n-1} - t_n)$. We assume that the heat flux at each of these time periods is constant. Thus, at the first time step,

$$\phi_{I} = -\frac{2\pi k_{e}}{W T_{D}(t_{D})} (T_{wb} - T_{ei})_{I}$$
(4.15)

Or,

$$T_{ei} - T_{wb,1} = \frac{W}{2\pi k_e} \phi_1 T_D(t_D)$$
(4.16)

The heat flow rate, ϕ_2 , during the second time step, $t_2 - t_1$, will be different from ϕ_1 . This situation will be represented by adding another constant heat source, which supply heat to the well at time > t_1 and whose magnitude is equal to $\phi_2 - \phi_1$. The

wellbore/formation interface temperature at this step, $T_{wb,2}$, is then the sum of the effects of these two heat sources and is given by,

$$T_{ei} - T_{wb,2} = \frac{W}{2\pi k_e} \left[\phi_i T_D(t_D) + (\phi_2 - \phi_i) T_D(t_D - t_{D,1}) \right]$$
(4.17)

Similarly, the third time period can be represented by three sources of heat supplying ϕ_1 from zero time, $\phi_2 - \phi_1$ since t_1 , and $\phi_3 - \phi_2$, since t_2 . Hence,

$$T_{ei} - T_{wb,3} = \frac{W}{2\pi k_e} \Big[\phi_1 T_D(t_D) + (\phi_2 - \phi_1) T_D(t_D - t_{D,1}) + (\phi_3 - \phi_2) T_D(t_D - t_{D,2}) \Big]$$
(4.18)

Hence for the nth time period,

$$T_{ei} - T_{wb,n} = \frac{W}{2\pi k_{e}} \sum_{n}$$
(4.19)

Where

$$\sum_{n} = \sum_{i=1}^{n} (\phi_{i} - \phi_{i-1}) T_{D}(t_{D} - t_{D,i-1})$$
(4.20)

and both φ_o and T_{D0} are zero.

The flowing wellbore fluid temperature is obtained from an energy balance between the wellbore fluid and the surrounding formation at the time of interest, t. The rate of heat transfer from the wellbore fluid to the wellbore/formation interface, in terms of the overall heat transfer coefficient for the wellbore, is given by,

$$\phi_n = -(2\pi r_{io} U_{io}) (T_f - T_{wb}) / W$$
(4.21)

Hence,

$$T_{wb,n} = T_{f_{\star}} + \frac{W \phi_n}{2 \pi r_{\mu} U_{\mu}}$$
(4.22)

Substituting this expression for $T_{wb,n}$ into Equation 4.19,

$$T_{ei} - T_{fn} = \frac{W \phi_n}{2 \pi r_{io} U_{io}} + \frac{W}{2 \pi k_e} \sum_{n=1}^{n-1} + \frac{W}{2 \pi k_e} (\phi_n - \phi_{n-1}) T_D(t_D - t_{D,n-1})$$
(4.23)

Where,

$$\sum_{n=1}^{n-1} = \sum_{i=1}^{n-1} (\phi_i - \phi_{i-i}) T_D(t_D - t_{D,i-1})$$
(4.24)

Or,

$$T_{ei} - T_{f,n} = \frac{\phi_n W}{2\pi} \left[\frac{1}{r_{io} U_{io}} + \frac{T_D(t_D - t_{D,n-1})}{k_e} \right] + \frac{W}{2\pi k_e} \sum_{n-1} - \frac{W}{2\pi k_e} (\phi_{n-1}) T_D(t_D - t_{D,n-1})$$
(4.25)

Or,

$$T_{ei} - T_{fn} = \frac{\phi_n W}{2\pi} \left[\frac{k_e + r_{io} U_{io} T_D (t_D - t_{D,n-1})}{k_e r_{io} U_{io}} \right] + \frac{W}{2\pi k_e} \sum_{n-1} - \frac{W}{2\pi k_e} (\phi_{n-1}) T_D (t_D - t_{D,n-1})$$
(4.26)

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Rewriting Equation 4.26 for ϕ_n ,

$$\Phi_{n} = \frac{C_{p}}{A_{n}} \left(T_{ei} - T_{f,n} \right) + T_{D} \left(t_{D} - t_{D,n-1} \right) \frac{\Phi_{n-1}}{B_{n}} - \frac{1}{B_{n}} \sum_{n-1}$$
(4.27)

Where,

$$A_{n} = \frac{W c_{p}}{2 \pi} \left[\frac{k_{e} + r_{to} U_{to} T_{D} (t_{D} - t_{D, n-1})}{k_{e} r_{to} U_{to}} \right]$$
(4.28)

And,

$$B_{n} = \frac{k_{e} + r_{io}U_{io}T_{D}(t_{D} - t_{D,n-1})}{r_{io}U_{io}}$$
(4.29)

Energy balance on the flowing fluid for a differential depth, dz, gives,

$$\frac{dT_f}{dz} = \frac{1}{c_p} \left[\phi_n - \frac{g \sin \theta}{g_c J} - \frac{V}{g_c J} \frac{dV}{dz} \right] + C_J \frac{dP}{dz}$$
(4.30)

Substituting the expression for φ_n from Equation 4.27 into Equation 4.30, we obtain,

$$\frac{dT_{f,n}}{dz} = \frac{T_{ei} - T_{f,n}}{A_n} - \frac{g \sin \theta}{g_c J c_p} - \frac{V}{g_c J c_p} \frac{dV}{dz} + C_J \frac{dP}{dz} + T_D (t_D - t_{D,n-1}) \frac{\Phi_{n-1}}{c_p B_n} - \frac{1}{c_p B_n} \sum_{n-1}$$
(4.31)

Or,

$$\frac{dT_{f,n}}{dz} = T_{ei} - \frac{T_{f,n}}{A_n} - \frac{g\sin\theta}{g_c J c_p} + \psi$$
(4.32)

Where,

$$\Psi = \sigma + T_D(t_D - t_{D,n-1}) \frac{\Phi_{n-1}}{c_p B_n} - \frac{1}{c_p B_n} \sum_{n-1}$$
(4.33)

And,

$$\sigma = C_{j} \frac{dP}{dz} - \frac{V(dV/dz)}{g_{c} J c_{p}}$$
(4.34)

The effect of various parameters on the term, σ (Equation 4.34), was investigated by Sagar et al. (1990) who proposed an empirical expression for its evaluation. This correlation is valid for flow rates less than 5 lb/sec and is shown below.

$$\phi = -0.00298 + 1.006 \times 10^{-6} P_{wh} + 1.91 \times 10^{-4} W - 1.05 \times 10^{-6} GOR$$

+ 3.229×10⁻⁵ API + 0.004 γ_{e} - 0.3551 g_{τ} (4.35)

Heat flow from the wellbore to the formation will vary with well depth. At the bottomhole, where the fluid temperature is the same as formation temperature, ϕ is zero. As the fluid moves upward in the well, the temperature difference between the wellbore fluid and the formation increases with consequent increase in the transfer of heat between the formation and the wellbore. Analytical expressions for wellbore fluid temperature as a function of well depth may be obtained from Equation 4.32 for two different assumptions about the variation of wellbore/formation interface heat fluxes, ϕ_n , as functions of well depth. It should be noted that we assume the geothermal gradient to be linear with depth, i.e,

$$T_{e_1} = T_{e_2b_1} - g_T z$$
 (4.36)

4.2.1.A Analytical Solution (Constant ψ)

Equation 4.32 can be rearranged as shown below as Equation 4.37,

$$\frac{dT_{f,n}}{dz} + \frac{T_{f,n}}{A_n} = \left[-\frac{g\sin\theta}{g_c J c_p} + \psi + \frac{T_{ei}}{A_n} \right] - \frac{g_T z}{A_n}$$
(4.37)

When heat fluxes are assumed not to vary significantly with depth, the terms in bracket on the right hand side of Equation 4.37 are constants while the last term is linear in the independent variable, z. Equation 4.37 is, therefore, a first order linear differential equation, which can be solved using integrating factor, I.F, given by,

$$I.F = e^{\int \frac{1}{A} dZ}$$

$$= e^{Z/A}$$
(4.38)

Multiplying each term by the I.F., we obtain,

$$\frac{d\left(T_{f,n}e^{Z/A}\right)}{dz} = \left[-\frac{g\sin\theta}{g_c J c_p} + \psi + \frac{T_{ei}}{A_n}\right]e^{Z/A} - \frac{g_T Z}{A_n}e^{Z/A}$$
(4.39)

To Integrate Equation 4.39 with respect to z, we note that the integral of $e^{z/A}$ is $Ae^{z/A}$ and that,

$$\frac{1}{A} \int e^{Z/A} g_T z \, dz = g_T z \, e^{Z/A} - \int g_T e^{Z/A} \, dz$$

$$= g_T z \, e^{Z/A} - A \, g_T e^{Z/A}$$
(4.40)

Hence the solution to Equation 4.37 is,

$$T_{f,n} e^{Z/A} = A e^{Z/A} \left[-\frac{g \sin \theta}{g_c J c_p} + \psi + \frac{T_{eibh}}{A} \right]$$

$$-g_T z e^{Z/A} + A g_T e^{Z/A} + IC$$

$$(4.41)$$

And,

$$T_{f,n} = A \left[-\frac{g \sin \theta}{g_c J c_p} + \psi + \frac{T_{eibh}}{A} \right] - g_T z + A g_T + IC e^{-Z/A}$$
(4.42)

where IC represents the integration constant. Noting that $T_{eibh} - zg_T$ equals formation temperature at the given depth, T_{ei} , we obtain,

$$T_{f.n} = T_{ei} + A \left[-\frac{g \sin \theta}{g_c J c_p} + \psi + g_T \right] + IC e^{-Z/A}$$
(4.43)

To evaluate the integration constant, IC, we use the condition that at the bottomhole (z=0) the fluid temperature is equal to the formation temperature ($T_{fn} = T_{eibh}$). Thus,

$$T_{fbh,n} = T_{eibh} + A \left[-\frac{g \sin \theta}{g_c J c_p} + \psi + g_7 \right] + IC$$
(4.44)

Or,

$$IC = T_{fbh,n} - T_{eibh} - A \left[-\frac{g \sin \theta}{g_e J c_p} + \psi + g_T \right] + IC$$
(4.45)

Hence,

$$T_{f,n} = T_{ei} + A_n \left(-\frac{g}{g_c} \frac{\sin\theta}{Jc_{pm}} + \psi + g_T \right) + e^{-Z/A} \left(T_{fbh} - T_{ebh} \right) - e^{-Z/A} A_n \left(-\frac{g}{g_c} \frac{\sin\theta}{Jc_{pm}} + \psi + g_T \right)$$

$$(4.46)$$

4.2.1.B Analytical Solution (Linear Variation of Heat Flux with depth)

Although the variation in heat flux with depth is small, it is possible to allow for this variation by using a linear variation of all ϕ 's with depth. Heat flux is zero at the bottomhole and attains a maximum negative value (heat flow is from the wellbore to the formation) at the wellhead. Thus, at any depth, z, heat fluxes, ϕ_i , are written (analogous to geothermal gradient) as,

$$\phi_i = F_i z \tag{4.47}$$

where F_i are positive numbers.

The parameter, ψ , given by Equation 4.33, is rewritten as,

$$\Psi = \sigma + T_D(t_D - t_{D,n-1}) \frac{z F_{n-1}}{c_p B_n} - \frac{z}{c_p B_n} \sum_{n-1}^{\infty} (4.48)$$

where,

$$\sum_{n=1}^{n} = \sum_{i=1}^{n-1} (F_i - F_{i-1}) T_D(t_D - t_{D,i-1})$$
(4.49)

Equation 4.37 then becomes,

$$\frac{dT_{f,n}}{dz} = -\frac{T_{f,n}}{A_n} + \frac{T_{eibh}}{A} - \frac{g\sin\theta}{g_c J c_p} + \sigma + \left[T_D(t_D - t_{D,n-1})\frac{F_{n-1}}{c_p B_n} - \frac{1}{c_p B_n}\sum_{n-1}^{r} - \frac{g_T}{A_n}\right]z$$
(4.50)

Equation 4.50, is very similar to Equation 4.37. The last term in Equation 4.50, which is linear in z, contains constants in addition to g_t . The IF remains the same and the solution is also very similar.

$$T_{f,n} = T_{eibh} - z F_T + A_n \left(-\frac{g}{g_c} \frac{\sin\theta}{Jc_{pm}} + \sigma + F_T \right) + e^{-Z/A} \left(T_{fbh} - T_{ebh} \right) - e^{-Z/A} A_n \left(-\frac{g}{g_c} \frac{\sin\theta}{Jc_{pm}} + \sigma + F_T \right)$$
(4.51)

Where,

$$F_{T} = g_{T} - T_{D}(t_{D} - t_{D,n-1}) \frac{A_{n}F_{n-1}}{B_{n}c_{p}} + \frac{A_{n}}{c_{p}B_{n}}\sum_{n-1}^{n} (4.52)$$

Here F_T is a function of n, but is independent of well depth.

4.3 Temperature Profile During Mud Circulation

Mud recirculation through tubings and tubing/casing annuli occurs during numerous operations, such as drilling, well kill operations, etc. The mud can enter either through the annulus or through the tube. If the mud enters through the annulus, then it returns through the tube and vice versa. The entering mud temperature at the wellhead is generally much lower than the bottomhole formation temperature. Thus, in flowing down through one channel and backing up through the other, the mud gains heat from the formation. The heat transfer rate for the mud in the annulus depends both on the formation temperature from which it gains heat, as well as on the tubing mud temperature to which it loses heat.

In the first two sections of this chapter we present two analytical solutions for the flowing mud temperature in the annulus and in the tubing for the two different system as a function of well depth. The solution is based on equating the heat loss from the formation to the heat gained by the mud in the tubing and in the annulus. The modeling approach utilizes the expression for the formation temperature distribution developed by Hasan and Kabir (1991). The resulting second order linear differential equation is solved in usual manner by adding the solution of the homogeneous equation to the particular solution for the inhomogeneous equation to obtain the final analytical expression.

One of the underlying assumptions of the above systems was the heat flux from the formation to the annular fluid remained constant throughout the entire operation. The effect of varying heat flux with operation time has been studied at the end of this chapter.



Figure 3. Circulating Fluid System

4.3.1 Mud Flow Down the Annuli and Up the Tubing

During a kill operation, mud is recirculated through tubing/casing annuli. In these and other cases, such as during drilling, it is necessary to estimate the flowing mud temperature in both the tubing and the annulus as a function of depth.

The entering mud temperature at the annulus wellhead, T_{si} , could be either slightly higher or slightly lower than the surface formation temperature. However, the formation temperature at the bottomhole is generally much higher than the annulus mud temperature. Thus, in flowing down the annulus and back up the tubing, the mud gains heat from the formation. The heat transfer rate for the mud in the annulus depends both on the formation temperature from which it generally gains heat, as well as on the tubing mud temperature to which it loses heat.

To obtain expressions for flowing mud temperature in the annulus and the drill tube, we set up an energy balance over a differential element of length, dz, of the annulus fluid as shown in Figure 4. Note that z is positive in the downward direction. Heat enters the element by convection at z, $q_a(z)$, and by conduction from the formation, q_F . Heat leaves the element by convection at z+dz, $q_a(z+dz)$ and by conduction and convection to the drill tube fluid, q_{u} . Thus,

$$q_a(z) - q_a(z + dz) = q_{ta} - q_F$$
 (4.53)

Or,

$$c_{P}\left[T_{a}(z) - T_{a}(z + dz)\right] = q_{Ia} - q_{F}$$
(4.54)

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 $q_{a}(z+dz)=c_{pm}T_{a}(z+dz)$



Figure 4. Schematic of Heat balance for Tubulars and Formation

Heat flow, q_F, from the formation to the wellbore is given by,

$$q_F = \frac{2\pi k_e}{WT_D} (T_{ei} - T_{wb}) dz$$
(4.55)

where q_F is heat transfer rate per unit length of the well per unit mass of mud. The dimensionless temperature, T_D , can be easily estimated from Equation 4.14 (Hasan and Kabir, 1991),

The wellbore/formation interface temperature, T_{wb} is related to the annulus mud temperature, T_a , by the overall heat transfer coefficient of the annulus system as follows,

$$q = \frac{2 \pi r_c U_a}{W} (T_{wb} - T_a)$$
(4.56)

The overall heat transfer coefficient, U_a , depends on the resistances to heat flow through the annulus mud, casing metal, and the cement, and is discussed in detail by Hasan and Kabir (1991).

Noting that the heat flowing from the formation to the annulus, q_F , equals q given by Equation 4.56, we eliminate T_{wb} from Equations 4.55 and 4.56 to we obtain the rate of heat transferred from the formation to the annulus,

$$q_F = \frac{2\pi}{W} \left[\frac{r_c U_a k_e}{k_e + T_D r_c U_a} \right] \left(T_{e_l} - T_a \right) dz$$
(4.57)

Or,

$$q_F = \frac{c_P}{A} \left(T_{e_I} - T_a \right) dz \tag{4.58}$$

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where,

$$A = \frac{c_{pm}W}{2\pi} \left[\frac{k_{e} + (r_{e}U_{a}T_{D})}{r_{e}U_{a}k_{e}} \right]$$
(4.59)

Heat transferred from the annulus mud to the mud in the drill tube is given by

$$q_{ta} = \frac{2 \pi r_{t} U_{t}}{W} (T_{a} - T_{t}) dz$$
(4.60)

Or,

$$q_{ta} = \frac{C_P}{B} \left(T_a - T_t \right) dz \tag{4.61}$$

Where,

$$B = \frac{W c_{P}}{2 \pi r_{L} U_{L}}$$
(4.62)

Hence Equation 4.54 is rewritten as,

$$c_{p}\left[T_{a}(z) - T_{a}(z + dz)\right] = \frac{c_{p}}{A}\left(T_{a} - T_{ei}\right)dz + \frac{c_{p}}{B}\left(T_{a} - T_{t}\right)dz \qquad (4.63)$$

Or,

$$A \frac{dT_a}{dz} = (T_{ei} - T_a) - (T_a - T_i) \frac{A}{B}$$
(4.64)

Equation 4.64 has two unknowns, the annulus mud temperature, T_a and the tubing mud temperature, T_i . An energy balance for the drill tube mud for the same differential element provides a second expression. Thus,
$$q_{i}(z+dz) - q_{a}(z) = -q_{ia}$$
(4.65)

Or,

$$c_{p}\left(T_{i}(z+dz) - T_{i}(z)\right) = c_{p} \frac{T_{i} - T_{a}}{B} dz$$
(4.66)

Or,

$$\frac{dT_{i}}{dz} = \frac{T_{i} - T_{a}}{B}$$
(4.67)

Or,

$$T_a = T_i - B \frac{dT_i}{dz}$$
(4.68)

Using Equation 4.68 to eliminate T_a from Equation 4.64, we obtain,

$$A \frac{dT_{t}}{dz} - AB \frac{dT_{t}^{2}}{d^{2}z} = T_{et} + B \frac{dT_{t}}{dz} - T_{t} + A \frac{dT_{t}}{dz}$$
(4.69)

Formation temperature, T_{ei}, is usually assumed to linearly increase with depth. Hence,

$$T_{1} = T_{2} + g_{\tau} z \tag{4.70}$$

And,

$$AB \frac{d^2 T_{t}}{dz^2} + B \frac{dT_{t}}{dz} - T_{t} + T_{ew} + g_T z = 0$$
(4.71)

Equation 4.71 is a second order linear differential equation with the following boundary conditions: $T_a = T_{ai}$ at the wellhead (i. e. when z = 0) and heat exchange between the tubing and wellbore mud is zero, i.e. $dT_t/dz = 0$, at the bottomhole (or $T_t=T_a$ when z=L). Equation 4.71, which is a inhomogeneous second order linear differential equation, is rewritten in the following form,

$$AB \frac{d^{2}T_{i}}{dz^{2}} + B \frac{dT_{i}}{dz} - T_{i} = -T_{ew} - g_{T}z \qquad (4.72)$$
$$= f(z)$$

 $f(z) = -T_{ew} - g_T z$

The particular solution of Equation 4.72 is easily obtained as

where,

$$T_{ip} = \mu z + \chi \tag{4.74}$$

(4.73)

Noting that the second derivative of T_{ψ} is zero, substitution of T_{ψ} as a solution to Equation 4.72 gives,

$$B\mu - \mu z - \chi = -T_{zz} - g_{z}$$
(4.75)

Equating coefficients of z and of the constants, we obtain,

$$\mu = g_T , \qquad \chi = T_{ew} + B g_T \qquad (4.76)$$

Thus the particular solution to Equation 4.72 is,

$$T_{tr} = g_T z + T_{tr} + B g_T \tag{4.77}$$

The complementary solution of Equation 4.72 is deduced from its homogeneous form (obtained by setting f(z) = 0),

$$AB \frac{d^2 T_i}{dz^2} + B \frac{dT_i}{dz} - T_i = 0$$
(4.78)

The characteristic polynomial equation for Equation 4.78 is

$$p(\lambda) = AB\lambda^2 - B\lambda - 1 = 0$$
 (4.79)

The solution to the quadratic equation gives the following two roots,

$$\lambda_{1} = -\frac{1}{2A} + \frac{1}{2A}\sqrt{1 + \frac{4A}{B}}$$
(4.80)

$$= -\frac{1}{2A} + \frac{1}{2A} \sqrt{1 + 4(r_c U_a T_D + k_e) \frac{r_i U_i}{r_c U_a k_e}}$$
(4.81)

Similarly,

$$\lambda_{2} = -\frac{1}{2A} - \frac{1}{2A} \sqrt{1 + 4(r_{c}U_{a}T_{D} + k_{e})\frac{r_{i}U_{i}}{r_{c}U_{a}k_{e}}}$$
(4.82)

The complementary solution to Equation 4.72 is then.

$$T_{\mu} = \alpha e^{\lambda_{1} Z} + \beta e^{\lambda_{2} Z}$$
(4.83)

and the complete solution, obtained by adding the particular and the complementary solutions, is

$$T_{t} = \alpha e^{\lambda_{1} Z} + \beta e^{\lambda_{2} Z} + g_{T} z + B g_{T} + T_{ew}$$
(4.84)

1

The expression for the annulus mud temperature can now be obtained using Equation 4.68 and Equation 4.84. Thus,

$$\frac{dT_{t}}{dz} = \alpha \lambda_{1} e^{\lambda_{1} z} + \beta \lambda_{2} e^{\lambda_{2} z} + g_{T}$$
(4.85)

And,

$$T_{a} = T_{t} - B \frac{dT_{t}}{dz}$$

$$= (1 - \lambda_{1}B) \alpha e^{\lambda Z} + (1 - \lambda_{2}B) \beta e^{\lambda_{2}Z} + g_{T}z + T_{ew}$$

$$(4.86)$$

The constants, α and β are obtained by applying the boundary conditions. Thus, at the wellhead, z = 0, $T_a = T_{ai}$, hence

$$T_{a} = T_{ai} = (1 - \lambda_{1}B)\alpha + (1 - \lambda_{2}B)\beta + T_{ew}$$

$$(4.87)$$

At the bottomhole (z = L), dT_1/dz is zero, i.e.,

$$\frac{dT_{t}}{dz}|_{L} = 0 = \alpha\lambda_{1}e^{\lambda_{1}L} + \beta\lambda_{2}e^{\lambda_{2}L} + g_{\tau}$$
(4.88)

Hence,

$$\alpha = -\frac{\beta \lambda_2 e^{\lambda_2 \ell} + g_T}{\lambda_1 e^{\lambda_1 \ell}}$$
(4.89)

Substituting this expression for α into Equation 4.87 and simplifying,

$$\beta = \frac{\left(T_{a_1} - T_{e_w}\right)\lambda_1 e^{\lambda_1 L} + g_T \left(1 - \lambda_1 B\right)}{\lambda_1 e^{\lambda_1 L} \left(1 - \lambda_2 B\right) - \lambda_2 e^{\lambda_2 L} \left(1 - \lambda_1 B\right)}$$
(4.90)

From Equation 4.89, then

$$\alpha = -\frac{\left(T_{ai} - T_{ew}\right)\lambda_2 e^{\lambda_2 L} + g_T \left(1 - \lambda_2 B\right)}{\lambda_1 e^{\lambda_1 L} \left(1 - \lambda_2 B\right) - \lambda_2 e^{\lambda_2 L} \left(1 - \lambda_1 B\right)}$$
(4.91)

4.3.2 Mud Flow Down the Tubing and Up the Annuli

When mud flows down the tubing and back up the annulus, the flow direction is reversed compared to the last case. Although the general approach for setting up the energy balance for the differential element remains the same, slight changes are needed in some of the expressions. We may still represent the heat flow from the formation to the annulus by Equation 4.58, and the heat flow from the tubing to the annulus by Equation 4.61, because in these cases the temperature difference driving force would appropriately account for the direction of heat flow. However, energy entering the mud in the differential element by convection is $q_a(z+dz)$ while that leaving is $q_a(z)$. Thus, Equations 4.53 and 4.54 are changed as follows,

$$q_{a}(z+dz) - q_{a}(z) = q_{ta} - q_{F}$$
 (4.92)

Or,

$$c_{p} \left[T_{a}(z + dz) - T_{a}(z) \right] = q_{ia} - q_{F}$$
(4.93)

Using Equations 4.58 and 4.61 Equation 4.93 is rewritten (similar to Equation 4.71) as,

$$A \frac{dT_a}{dz} = (T_a - T_i) \frac{A}{B} - (T_{ei} - T_a)$$

$$(4.94)$$

Energy balance for the mud in the drill tube leads to the following equation, which has the opposite sign to that of Equation 4.65,

$$q_{1}(z) - q_{2}(z + dz) = -q_{12}$$
(4.95)

Or,

$$T_a = T_i + B \frac{dT_i}{dz}$$
(4.96)

Combining Equations 4.94 and 4.96 to eliminate $T_{\scriptscriptstyle a}$, we obtain,

$$AB \frac{d^2 T_{t}}{dz^2} - B \frac{dT_{t}}{dz} - T_{t} + T_{ew} + g_T z = 0$$
(4.97)

Equation 4.97 can be solved in a manner similar to Equation 4.91 with the following boundary conditions: $T_t = T_u$ at the wellhead (i. e. when z = 0) and $dT_t/dz = 0$ at the bottomhole (i.e. when z = L). The solution is,

$$T_{t} = \gamma e^{\xi_{1}Z} + \delta e^{\xi_{2}Z} + g_{T}Z - B g_{T} + T_{ew}$$
(4.98)

And,

$$T_{a} = (1 + \xi_{1}B) \gamma e^{\xi_{1}Z} + (1 + \xi_{2}B) \delta e^{\xi_{2}Z} + g_{\tau}z + T_{ew}$$
(4.99)

Where,

$$\gamma = -\frac{\left(T_{ii} + Bg_T - T_{ew}\right)\xi_2 e^{\xi_2 L} + g_T}{\xi_1 e^{\xi_1 L} - \xi_2 e^{\xi_2 L}}$$
(4.100)

$$\delta = \frac{\left(T_{ii} + B g_T - T_{ew}\right) \xi_1 e^{\xi_1 L} + g_T}{\xi_1 e^{\xi_1 L} - \xi_2 e^{\xi_2 L}}$$
(4.101)

$$\xi_{1} = \frac{1}{2A} + \frac{1}{2A} \sqrt{1 + 4(r_{c}U_{a}T_{D} + k_{e}) \frac{r_{i}U_{i}}{r_{c}U_{a}k_{e}}}$$
(4.102)

$$\xi_{2} = \frac{1}{2A} - \frac{1}{2A} \sqrt{1 + 4(r_{c}U_{a}T_{D} + k_{e}) \frac{r_{i}U_{i}}{r_{c}U_{a}k_{e}}}$$
(4.103)

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4.3.3 Effect of Varying Heat Flux During Mud Circulation

We pointed out earlier that the heat flux from the formation to the annular fluid changes with the operation time. The concept of varying heat flux can be applied to the mud circulation system using the same principle that was earlier used in the single pipe system. Heat flux at any time is a function of previous heat flux and can be expressed as,

$$\Phi_{n} = \frac{C_{p}}{A_{n}} (T_{ei} - t_{a,n}) + T_{D} (t_{D,n} - t_{D,n-1}) \frac{\Phi_{n-1}}{B_{n}} - \frac{1}{B_{n}} \sum_{i=1}^{n-1} (\Phi_{i} - \Phi_{i-1}) T_{D} (t_{D,n} - t_{D,i-1}) \quad .$$
(4.104)

where,

$$A_{n} = \frac{W c_{p}}{2\pi} \left[\frac{k_{e} + r_{c} U_{a} T_{D} (t_{D,n} - t_{D,n-1})}{k_{e} r_{c} U_{a}} \right]$$
(4.105)

and

$$B_{n} = \frac{k_{e} + r_{c} U_{a} T_{D} (t_{D,n} - t_{D,n-1})}{r_{c} U_{a}}$$
(4.106)

Heat transferred from the annulus mud to the mud in the tube is given by,

$$\phi_{ia} = \frac{2 \pi r_i U_i}{W} (T_a - T_i)$$

$$= \frac{c_p}{E} (T_a - T_i)$$
where, $E = \frac{W c_p}{2 \pi r_i U_i}$
(4.107)

The final form of the energy balance over the differential length, dz of the annulus fluid includes two new terms to account for the varying heat flux.

$$c_{\rho} \left[T_{a}(z) - T_{a}(z + dz) \right] = (\phi_{ia} - \phi) dz$$
(4.108)

Or,

$$A_{n}\frac{dT_{a}}{dz} = (T_{ei} - T_{a,n}) - \frac{A_{n}}{E}(T_{a} - T_{i}) + \frac{A_{n}\phi_{n-1}}{B_{n}c_{p}}T_{D}(t_{D,n} - t_{D,n-1}) - \frac{A_{n}}{B_{n}c_{p}}\sum_{i=1}^{n-1} (\phi_{i} - \phi_{i-1}) T_{D}(t_{D,n} - t_{D,i-1})$$

$$(4.109)$$

The above equation has two unknowns, the annulus mud temperature, T_a and the tubing mud temperature, T_t . Earlier it was shown that energy balance over the differential length on the tube side provides the following expression.

$$T_a = T_t - E \frac{dT_t}{dz}$$
(4.110)

Using the above equation to eliminate T_a from Equation 4.109, we obtain,

$$A_{n}E \frac{d^{2}T_{t}}{dz^{2}} + T_{ew} + g_{t}z - T_{t} + E \frac{dT_{t}}{dz} + \frac{A_{n}\phi_{n-1}}{B_{n}c_{p}}T_{D}(t_{D,n} - t_{D,n-1}) - \frac{A_{n}}{c_{p}B_{n}}\sum_{i=1}^{n-1} (\phi_{i} - \phi_{i-1}) T_{D}(t_{D,n} - t_{D,i-1}) = 0$$
(4.111)

This equation is a second order linear differential equation with boundary conditions : at the wellhead, $Ta = T_{ai}$ and at the bottomhole, heat exchange between the tubing and annulus is zero, i.e. $dT_{ai}/dz = 0$. Equation 4.111, which is a inhomogeneous second order linear differential equation, can be rewritten in the following form,

$$A_{n}E\frac{d^{2}T_{t}}{dz^{2}} + E\frac{dT_{t}}{dz} - T_{t} = -T_{ew} - g_{t}z - \Omega_{1} + \Omega_{2}$$
(4.112)

where,

$$\Omega_{1} = \frac{A_{n} \Phi_{n-1}}{B_{n} c_{p}} T_{D} (t_{D,n} - t_{D,n-1})$$

$$\Omega_{2} = \frac{A_{n}}{c_{p} B_{n}} \sum_{i=1}^{n-1} (\Phi_{i} - \Phi_{i-1}) T_{D} (t_{D,n} - t_{D,i-1})$$
(4.113)

The particular solution of Equation 4.112 can easily be obtained as,

$$T_{tr} = g_{t}^{2} + E g_{t} + T_{ew} + \Omega_{1} - \Omega_{2}$$
(4.114)

and the complementary solution of the Equation can be deduced from its homogeneous form,

$$A_{n}E\frac{d^{2}T_{i}}{dz^{2}} + E\frac{dT_{i}}{dz} - T_{i} = 0$$
(4.115)

and has been found to be,

$$T_{\mu} = \alpha e^{\lambda_1 Z} + \beta e^{\lambda_2 Z}$$
(4.116)

where α , β , λ_1 and λ_2 are constants. The solution of the quadratic equation (Equation 4.115) gives the value of λ_1 and λ_2 , whereas values of α and β can be obtained by applying the boundary conditions. The complete solution obtained by adding the particular and complementary solution is as follows,

$$T_{t} = \alpha e^{\lambda_{1} Z} + \beta e^{\lambda_{2} Z} + g_{t} Z + E g_{t} + T_{ew} + \Omega_{1} - \Omega_{2}$$
(4.117)

and the mud temperature in the annulus,

$$T_{a} = (1 - \lambda_{1}E) \alpha e^{\lambda_{1}Z} + (1 - \lambda_{2}E) \beta e^{\lambda_{2}Z} + g_{1}z + T_{ew} + \Omega_{1} - \Omega_{2}$$
(4.118)

By applying the boundary conditions the following values of α and β were obtained,

$$\alpha = - \frac{(T_{ai} - T_{ew} - \Omega_1 + \Omega_2)\lambda_2 e^{\lambda_2 L} + g_r(1 - \lambda_2 E)}{\lambda_1 e^{\lambda_1 L} (1 - \lambda_2 E) - \lambda_2 e^{\lambda_2 L} (1 - \lambda_1 E)}$$
(4.119)

$$\beta = \frac{(T_{ai} - T_{ew} - \Omega_1 + \Omega_2)\lambda_1 e^{\lambda_1 L} + g_i(1 - \lambda_1 E)}{\lambda_1 e^{\lambda_1 L} (1 - \lambda_2 E) - \lambda_2 e^{\lambda_2 L} (1 - \lambda_1 E)}$$
(4.120)

Down the Tube Up the Annulus :

When mud flows down through the tubing and backs up through the annulus, the flow direction is reversed compared to the last case. Although general approach for setting up the energy balance for the differential element remains the same, slight variation is observed in the expression,

$$c_{p}\left[T_{a}(z + dz) - T_{a}(z)\right] = (\phi_{ta} - \phi) dz$$
(4.121)

Substituting the values of φ_{u} and $\varphi,$ we obtain

$$A_{n}\frac{dT_{a}}{dz} = -(T_{ei} - T_{a,n}) + \frac{A_{n}}{E}(T_{a} - T_{i}) - \frac{A_{n}\phi_{n-1}}{B_{n}c_{p}}T_{D}(t_{D,n} - t_{D,n-1}) + \frac{A_{n}}{B_{n}c_{p}}\sum_{i=1}^{n-1}(\phi_{i} - \phi_{i-1})T_{D}(t_{D,n} - t_{D,i-1})$$

$$(4.122)$$

Energy balance in the tube leads the following equation,

$$T_a = T_i + E \frac{dT_i}{dz}$$
(4.123)

Using the above equation to eliminate T_a from Equation 4.122, we get,

$$A_{n}E\frac{d^{2}T_{i}}{dz^{2}} + T_{ew} + g_{i}z - T_{i} - E\frac{dT_{i}}{dz} + \frac{A_{n}\phi_{n-1}}{B_{n}C_{p}}T_{D}(t_{D,n} - t_{D,n-1}) - \frac{A_{n}}{C_{p}B_{n}}\sum_{i=1}^{n-1} (\phi_{i} - \phi_{i-1})T_{D}(t_{D,n} - t_{D,i-1}) = 0$$

$$(4.124)$$

This equation is a second order linear differential equation with boundary conditions : at the wellhead, $T_t = T_{ti}$ and at the bottomhole, heat exchange between the tubing and annulus is zero, i.e. $dT_t/dz = 0$. Equation 4.124, which is an inhomogeneous second order linear differential equation, can be rewritten in the following form,

$$A_{n}E\frac{d^{2}T_{t}}{dz^{2}} - E\frac{dT_{t}}{dz} - T_{t} = -T_{ew} - g_{t}z - \Omega_{1} + \Omega_{2}$$
(4.125)

where Ω_1 and Ω_2 are the same constants described earlier. The particular solution of Equation 4.125 can easily be obtained as.

$$T_{tr} = g_{t} z + E g_{t} + T_{rw} + \Omega_{1} - \Omega_{2}$$
(4.126)

and the complementary solution of the Equation can be deduced from its homogeneous form

$$A_{n}E\frac{d^{2}T_{i}}{dz^{2}} - E\frac{dT_{i}}{dz} - T_{i} = 0$$
(4.127)

and has been found to be,

$$T_{1} = \gamma e^{\frac{5}{2}} + \delta e^{\frac{5}{2}}$$
(4.128)

where γ , δ , ξ_1 and ξ_2 are constants. The solution of the quadratic equation (Equation 4.127) gives the value of ξ_1 and ξ_2 , whereas values of γ and δ can be obtained by applying the boundary conditions.

The complete solution obtained by adding the particular and complementary solution is as follows

$$T_{t} = \gamma e^{\xi_{t} Z} + \delta e^{\xi_{t} Z} + g_{t} z - E g_{t} + T_{ew} + \Omega_{1} - \Omega_{2}$$
(4.129)

and the mud temperature in the annulus,

$$T_{a} = (1 + \xi_{1}E) \gamma e^{\xi_{1}Z} + (1 + \xi_{2}E) \delta e^{\xi_{2}Z} + g_{1}Z + T_{ew} + \Omega_{1} - \Omega_{2}$$
(4.130)

By applying the boundary conditions the following values of α and β were obtained,

$$\gamma = - \frac{(T_{ii} + E g_i - T_{ew} - \Omega_1 + \Omega_2)\xi_2 e^{\xi_2 L} + g_i}{\xi_1 e^{\xi_1 L} - \xi_2 e^{\xi_2 L}}$$
(4.131)

$$\delta = \frac{(T_{ii} + Eg_i - T_{ew} - \Omega_1 + \Omega_2)\xi_1 e^{\xi_1 t} + g_i}{\xi_1 e^{\xi_1 t} - \xi_2 e^{\xi_2 t}}$$
(4.132)

CHAPTER 5

RESULTS AND DISCUSSION

One of the objective of this work is to develop a unified model for void fraction and pressure gradient in two-phase flow. This model is then used to develop a simulator that is uniquely useful to the petroleum industry. The other objective is to study the complex heat transfer problems encountered in wellbores. This includes prediction of temperature profile during production, injection, and mud circulation. In the first section of this chapter we discuss some of the two-phase simulation results. In the second section some of the important aspects of heat transfer is presented.

5.1 Pressure and Void Fraction Profile

Two approaches may be taken to calculate the pressure profile in an wellbore. In both the approaches, the calculations start out with the wellhead or the bottom condition whichever is known to the user. In one method, the pressure gradient is determined at the known condition and the gradient is multiplied by a certain length (usually a small percentage of the total length, i.e., 1%), which gives pressure at the end of this length. The physical properties of the system, including the superficial velocities are calculated at the new point and a new gradient is calculated. The procedure is repeated until the entire well is traversed in this manner. The drawback of this procedure is that the pressure gradient calculated at the beginning of each section is different from the average gradient for the section. The gradient at any point is dependent on fluid properties which are function of system pressure. So, we use Runge Kutta method to better this prediction. The calculation procedure of our program, which uses the second approach, is somewhat different than most of the available programs. We considered pressure as the independent variable, and the dependent variable length is calculated using the pressure gradient. The advantage of taking pressure as the independent variable is that any error in pressure gradient estimation does not directly influence the fluid property and the gradient calculation. Better prediction is expected from this procedure. It is possible to devise other, numerically more efficient procedure, for pressure prediction in an oilwell. However, property correlations are not very accurate and gradient change is not steep enough, for any further sophistication of the numerical procedure.

The pressure and void fraction profile of two different systems as calculated by the models described in the earlier chapters are shown in Figures 5 and 6. The description of the systems are given in Tables 1 and 2 respectively. Figure 5 shows the pressure and void fraction profile for 4355 ft vertical well producing 855 bbl/day of crude. The bottomhole condition is known in this case and simulation progresses from bottom to top. Besides bubbly and slug, single phase flow is observed in this case. Single phase flow persists from the bottomhole to a depth of around 3200 ft. The pressure within this region is high enough for the gas phase to be dissolved in the liquid phase. As the crude passes this depth, the lower pressure causes gas to come out of the liquid phase forming a distinct phase and bubbly flow starts. Bubbly flow continues up to a



Figure 5. Pressure (and Void Fraction) vs. Depth in Vertical Upward Flow in a Pipe

depth of 1030 ft and transition to slug flow occurs at this depth. An abrupt lowering of void fraction is observed when the pattern changes from bubbly to slug flow. This abrupt change is due to the higher velocity of Taylor bubbles compared to the rise velocity of smaller bubbles that cause the gas phase to move faster and void fraction to decrease.

Figure 6 corresponds to flow in 4450 ft annulus inclined 80° to the horizontal. The flow is in downward direction and the surface condition is known in this case. Simulation progresses from top to bottom. Bubbly flow persists from the wellhead up to a length of about 2150 ft. The conditions are such that slug flow is not observed in this case. At the depth of 2150 ft, the pressure becomes high enough for the gas phase to get dissolved in the liquid phase. Single phase flow starts at that point and it continues up to the bottomhole.

TABLE 1Wellbore and Fluid Data(Upward Flow in Vertical Pipe)		
Well Depth, ft	4355	
Production Rate, STB/day	855	
Tube Diameter, in	0.249	
Bottomhole Pressure, psia	1715	
Bottomhole Temperature, F	153	
Gas to Oil Ratio	185	
Specific Gravity of Gas	0.75	
Specific Gravity of Liquid	0.98	
Angle of inclination with Horizontal, deg	90	
Surface Tension, lb/sec ²	0.058	



Figure 6. Pressure (and Void Fraction) vs. Depth in Inclined Downward Flow in Annulus

TABLE 2Wellbore and Fluid Data(Downward Flow in Inclined Annulus)		
Well Depth, ft	4475	
Production Rate, STB/day	855	
Tube Diameter, in	0.249	
Annuli Diameter, in	0.416	
Surface Pressure, psia	1715	
Surface Temperature, F	153	
Gas to Oil Ratio	185	
Specific Gravity of Gas	0.75	
Specific Gravity of Liquid	0.98	
Angle of inclination with Horizontal, deg	80	
Surface Tension, lb/sec ²	0.058	

5.2 Effect of Varying Heat Flux during Production

In the second section of Chapter 3, we observed that the present approach of estimating fluid temperature during production and injection assumes heat flux to remain constant throughout the production time. However, it is intuitively obvious that heat flux decreases with the production time. We developed two expressions in chapter 4 for estimating fluid temperature in wellbores which account for the variation of heat flux with time. The first expression assumes that the parameter, ψ does not vary significantly with well depth. In reality, however, ψ is a complex function of heat flux, time, and pressure gradient. In deed, the heat flux is zero at the bottomhole (because fluid temperature and

formation temperature are same) and maximum at the wellhead and varies throughout the well depth. In the second approach, we make the more realistic assumption of linear variation of heat flux with depth.

A number of simulations has been run for an oilwell parameters for which are given in Table 3. Besides using the two approaches explained above, numerical solution of the governing differential equation was also sought. Five different solutions with 1, 5, 10, 50, and 100 time steps (= n) were examined for each approach. The number of time step n = 100 signifies that the total production time of 158 hours were divided in 100 equal intervals, and for each 1.58 hour period, the heat flux was assumed to remain at a constant value. Here, n = 1 signifies that heat flux remains constant throughout the entire production time of 158 hours and as such coincides with the solution presently available in the literature. It has been found that although there is significant variation between the results of single step and 100 steps, the results of 50 and 100 time step simulations are close. The simulation with more discritization (such as n = 150) resulted in almost the same profile. So, we concluded that 100 time steps are sufficient for this case. The temperature and heat flux profiles obtained by the simulations are discussed in the following paragraphs. All the profiles will represent the end of production time profile unless otherwise noted.

A number of scenarios have been examined and the results are presented in two different sets of figures. The first set accounts convection in tubing/casing annulus in determining the overall heat transfer coefficient, while the second set does not. The temperature and heat flux profiles with constant ψ assumption are shown in Figures 7

TABLE 3Wellbore and Fluid Data During Production in Wellbore		
Well Depth , ft Production Rate, lb/hr Tube Diameter, in Casing Diameter, in Wellbore Diameter, in Specific Gravity of Crude, API Formation Thermal Conductivity, Btu/hr ft F Cement Thermal Conductivity, Btu/hr ft F	5400 8856 2.875 7.0 9.0 34.3 0.83 4.021	
Specific Heat of fluid, Btu/lb F Surface Earth Temperature, F Geothermal Gradient, Production Time, hr Thermal Diffusivity, ft/hr ²	0.947 76 0.005926 158 0.04	

and 8. The results show significant differences in temperature and heat flux profile as we incorporate the concept of varying heat flux with time. With this approach (ψ =constant), heat flux at any given depth at the end of production time has been found to increase as we discritize the production time. But the physical system suggests that heat flux should decrease if the production time is discritized. This discrepancy suggests that the assumption of constant ψ was probably not a very good one.

The temperature and heat flux profiles with the assumption of linear variation of heat flux with depth are shown in Figures 9 and 10. These simulations also accounted for convection. The solutions have been carried out under the same conditions as the earlier case and five different scenarios at the same number of time steps are presented.

Temperature Profile in the Wellbore Constant Psi, With Convection



Figure 7. Temperature vs. Depth in a 5400 ft Wellbore (Constant ψ and With Convection)



Figure 8. Heat Flux vs. Depth in a 5400 ft Wellbore (Constant ψ and With Convection)

Temperature Profile in the Wellbore Linear Phi, With Convection



Figure 9. Temperature vs. Depth in a 5400 ft Wellbore (Linear Heat Flux and With Convection)

Heat Flux Profile in the Wellbore Linear Phi, With Convection



Figure 10. Heat Flux vs. Depth in a 5400 ft Wellbore (Linear Heat Flux and With Convection)



Figure 11. Temperature vs. Depth in a 5400 ft Wellbore (Numerical Solution)

Heat Flux Profile in the Wellbore Numerical Solution



Figure 12. Heat Flux vs. Depth in a 5400 ft Wellbore (Numerical Solution)

The results show a difference in profile with discritization of production time. But the extent of variation is less compared to the earlier case of constant ψ . The heat flux profiles show that as the total production time is discritized, the heat flux decreases, which is consistent with the physical system. The numerical solutions of the governing differential equation are presented in Figures 11 and 12. The nature of this problem does not easily allow convection to be taken into account in the numerical procedure. The conductivity of formation was changed from 0.83 to 1.40 in order to compensate for convection in the overall heat transfer coefficient term. The numerical solution matches with the solutions obtained from linear variation of heat flux approach. The consistency with the physical system and close match with the numerical results suggest that the linear variation of heat flux with depth assumption is a reasonable one.

The second set of results are shown in Figures 13 to 18. These results do not account for convection in determining the overall heat transfer coefficient. Figure 13 and 14 show the temperature and pressure profile obtained by using the constant ψ assumption. Similarly Figures 15 and 16 show the solution that resulted from the assumption of linear variation of heat flux and Figures 17 and 18 are from numerical solutions. Same conclusion regarding the validity of the linear heat flux assumption can be drawn from this set of results as well. A number of other simulations have been run for a 8000 ft oilwell. The results of these simulations, along with the description of the system, are presented in Appendix C. These results also support the conclusion that linear variation of heat flux assumption is a reasonable one.

Temperature Profile in the Wellbore Constant Psi, Without Convection



Figure 13. Temperature vs. Depth in a 5400 ft Wellbore (Constant ψ and Without Convection)

Heat Flux Profile in the Wellbore Constant Psi, Without Convection



Figure 14. Heat Flux vs. Depth in a 5400 ft Wellbore (Constant ψ and Without Convection)



Temperature Profile in the Wellbore Linear Phi, Without Convection

Figure 15. Temperature vs. Depth in a 5400 ft Wellbore (Linear Heat Flux and Without Convection)

Heat Flux Profile in the Wellbore Linear Phi, Without Convection



Figure 16. Heat Flux vs. Depth in a 5400 ft Wellbore (Linear Heat Flux and Without Convection)



Figure 17. Temperature vs. Depth in a 5400 ft Wellbore (Numerical Solution)



Figure 18. Heat Flux vs. Depth in a 5400 ft Wellbore (Numerical Solution)

5.3 Heat Transfer during Mud Circulation

The knowledge of accurate temperature with time during mud circulation is of prime importance in many operations including drilling, cementing, etc. In the third part of chapter 4, we developed two sets of expressions to estimate circulating fluid temperature in tube and annuli. The first set assumes that heat flux between the annulus fluid and the formation remains constant throughout the whole production time, while the second set accounts for the variation of heat flux with time.

TABLE 4Wellbore and Fluid Data during Mud Circulation		
Well Depth, ft	15000.0	
Drillstem OD, ft	0.552	
Drillbit Size, ft	0.698	
Circulation Rate, bbl/hr	300.0	
Mud Density, lb/gal	10.0	
Formation Density, lb/cu.ft	165.0	
Geothermal Gradient	0.0127	
Mud Specific Heat, Btu/lb F	0.4	
Formation Specific Heat, Btu/lb F	0.2	
Mud Viscosity, lb/ft.hr	110.0	
Formation Thermal Conductivity, Btu/lb-F	1.3	
Mud thermal Conductivity, Btu/lb-F	1.0	
Inlet Mud Temperature, F	75.0	
Surface Earth Temperature, F	59.5	

Both the approaches have been used to estimate the tube and annuli fluid temperatures in a well and the well's description is available in Table 4. The simulations



Temperature Profile in Mud Circulation Down the Annuli and Up the Tube

Figure 19. Temperature vs. Depth During Mud Circulation (Down the Annulus, Up the Tube)

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Temperature Profile in Mud Circulation Down the Tube and Up the Annuli



Figure 20. Temperature vs. Depth During Mud Circulation (Down the Tube, Up the Annulus)

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were carried out under two different circumstances. In the first case, fluid was assumed to flow down the annulus, and up the tubing, whereas in the second case the flow direction was reversed. While calculating fluid temperature using the varying heat flux concept, the total circulation time was discritized into 100 equal intervals so that convergence could be obtained. The resulting temperature profiles are shown in Figures 19 and 20. A significant difference in temperature profile is observed with the incorporation of varying heat flux concept.

CHAPTER 6

CONCLUSION AND RECOMMENDATION

Conclusions

• A unified model for two-phase flow was developed using the models presently available in the literature. This model was then used to develop a simulator which is useful to the petroleum industry. The simulator first determines the existing flow pattern from flow rates and fluid properties. Then void fraction and pressure gradient were estimated for that pattern. Pressure was taken as the independent variable, and length is calculated for a certain change in pressure. Forward marching technique is used to estimate the length for subsequent pressure change until the total well length is traversed.

• An expression for fluid temperature during production and injection as a function of well depth and operation time was developed from an energy balance between the fluid and the formation. Particular emphasis was given to the appropriate boundary condition in order to account for the variation of heat flux between the wellbore and formation. Significant difference is observerd between the results of the proposed solution and the presently available solution.

• Expressions for fluid temperature in tube and annulus during mud circulation as a function of well depth and circulation time have been developed. Both the cases - flow down the annulus, up the tube; and flow down the tube, up the annulus, were modeled. The use of superposition principle affects the temperature profiles significantly.

Recommendations

• As the flow pattern changes from bubbly to slug, or slug to churn an abrupt lowering of void fraction is observed. This is due to the higher velocity of Taylor bubbles compared to the smaller bubbles. However, in reality a more gradual transition is expected. We recommend some modifications in the model so that a gradual transition can be accounted for.

• The developed model is applicable only at steady state flow. In practice, under many circumstances, the system does not attain steady state. This type of situations are very common in well testing. Not much work has been done and considerable oppourtunity lies in this area.

• We recommend that the concept of varying heat flux be used in the temperature estimation process during the gas lift operation and also in the pressure transient analysis. The transient heat transfer model presented here could be coupled with the momentum balance in the pressure transient analysis. This heat transfer model would also be useful in various situations where temperature plays an important role, such as in the analysis of transient physical response of the wellbore fluid during phase redistribution.

APPENDICES

APPENDIX A

NOMENCLATURE

- A Inverse relaxation distance, ft (m).
- A_n Inverse relaxation distance for nth time step, ft(rn)
- B Constant used in Equation 4.61, defined by Equation 4.62,
- c_a Heat capacity of annulus fluid, Btu/lb °F (kJ/kg °C).
- c_p Heat capacity of wellbore fluid, Btu/lb °F (kJ/kg °C).
- c. Heat capacity of earth, Btu/lb °F (kJ/kg °C).
- C_J Joule-Thompson coefficient, dimensionless.
- C_o Flow parameter in bubbly flow, dimensionless.
- C₁ Flow parameter in slug flow.dimensionless
- D Pipe diameter, ft (m).
- D_i Diameter of inside pipe, ft (m)
- D_0 Diameter of outside pipe, ft (m)
- E Entrainment factor, dimensionless
- E_g Gas void fraction, dimensionless
- E₁ Liquid Volume Fraction, dimensionless
- f Friction factor, dimensionless.
- f_f Film friction factor in annular flow, dimensionless.
- f_g Gas void fraction, or in-situ volume fraction, dimensionless.
- f_m Friction factor for two-phase flow, dimensionless.
- g Acceleration due to gravity, ft/sec^2 (m/s²).
- g_c Conversion factor, 32.2 $lb_m ft/lb_f s^2$, unity in SI units, dimensionless.
- g_T Geothermal temperature gradient, °F/ft (°C/m).

- Gr. Grashof number, defined by Equation 52, dimensionless.
- h Heat transfer coefficient, Btu/°F sec ft (kJ/°F s m)
- h_c Convective heat transfer coefficient for annulus fluid, Btu/°F s ft (kJ/°C s m)
- h, Radiative heat transfer coefficient for the annulus, Btu/°F sec ft (kJ/°F s m)
- h_t Forced convection heat transfer coefficient for the tubing fluid, Btu/°F sec ft (kJ/°C s m)
- H Fluid enthalpy, Btu/lb (kJ/kg).
- I Definite integral given by Equation 3.23, dimensionless.
- J Conversion Factor, 778 ft lb_f/Btu.
- J₀ Zero-order Bessel function of the first kind, dimensionless.
- J₁ First-order Bessel function of the first kind, dimensionless.
- k Conductivity, Btu/ft °F (kJ/m °C).
- k, Conductivity of the annulus fluid Btu/ft °F (kJ/m °C).
- k_c Conductivity of the casing material, Btu/ft °F (kJ/m °C).
- k_{cem} Cement conductivity, Btu/ft °F (kJ/m °C).
- k_e Earth (formation) conductivity, Btu/ft °F (kJ/m °C).
- k_{ins} Conductivity of the insulating material, Btu/ft °F (kJ/m °C).
- k_t Conductivity of the tubing material, Btu/ft °F (kJ/m °C).
- L Length of the well, ft(m)
- L_c Length of a *cell* in the cellular model of slug flow, ft (m).
- L_s Length of the liquid slug in a *cell* in slug flow, ft (m).
- L_T Length of the Taylor bubble portion in a *cell* in slug flow, ft (m).
- P Wellbore fluid pressure, psi (kPa).
- Pr Prantl number, defined by Equation 53, dimensionless.
- q Heat flow rate from, or to, the wellbore, Btu/hr (kJ/hr).
- q_F Heat flow between the formation and wellbore, Btu/hr (KJ/hr)
- q_{ta} Heat flow between the tube and annulus, Btu/hr (KJ/hr)
- r Radial distance from the wellbore, ft (m).
- r_{ci} Inside radius of the casing ft (m).
- r_{co} Outside radius of the casing ft (m).

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- r_{u} Inside radius of the tubing ft (m).
- r_{10} Outside radius of the tubing ft (m).
- r_{wb} Outside radius of the wellbore (or cement) ft (m).
- t Time, sec (s).
- t_D Dimensionless time = $\alpha t/r_{wb}$, dimensionless.
- T Temperature, °F (°C).
- T_{ci} Casing inside surface temperature, °F (°C).
- T_{co} Casing outside surface temperature, °F (°C).
- $T_{\rm D}$ Dimensionless temperature = $(2\pi k_{\rm e})(T_{\rm wb} T_{\rm ei})/(W\phi)$, dimensionless.
- T_e Earth temperature at any given depth and radial distance from well, °F (°C).
- T_{ei} Earth temperature at any given depth and far away from the well, °F (°C).
- T_{eibh} Earth temperature at the bottomhole and far away from the well, °F (°C).
- T_{ewh} Earth temperature at the wellhead and far away from the well, °F (°C).
- T_f Wellbore (tubing) fluid temperature, °F (°C).
- T_{ins} Insulation (outside) surface temperature, °F (°C).
- T_{wb} Wellbore/earth interface temperature, °F (°C).
- u Dummy variable for integration in Eqs., dimensionless.
- V Fluid velocity, ft/sec (m/s).
- V_g In-situ gas velocity, ft/sec (m/s).
- V_{sg} Superficial gas velocity, ft/sec (m/s).
- V_{sl} Superficial liquid velocity, ft/sec (m/s).
- V_m Two-phase mixture velocity, ft/sec (m/s).
- V_t Terminal rise velocity of small bubbles, ft/sec (m/s).
- V_{tr} Terminal rise velocity of Taylor bubbles, ft/sec (m/s).
- W Total mass flow rate, lb_m/sec (kg/s).
- x Quality, mass fraction of the gas phase, dimensionless.
- z Variable well depth from surface, ft (m).
- z_{bh} Total well depth from surface, ft (m).

Greek Letters

- β Constant used in the solution of differential equation during mud circulation
 γ Constant used in the solution of differential equation during mud circulation
 δ Constant used in the solution of differential equation during mud circulation
 θ Pipe inclination angle with horizontal, degree.
- μ Fluid viscosity, cp.
- ρ Density, lb_m/ft^3 (kg/m³)
- ρ_e Earth density, lb_m/ft^3 (kg/m³)
- ρ_{g} Gas density, lb_{m}/ft^{3} (kg/m³)
- ρ_1 Liquid density, lb_m/ft^3 (kg/m³)
- ρ_m Two-phase mixture density, lb_m/ft^3 (kg/m³)
- σ Surface tension, lb_m/sec^2 (N/m).
- λ Constant used in the solution of differential equation during mud circulation
- ϕ Heat flux between formation and wellbore, Btu/ft²hr (KJ/m²hr)
- χ Constant used in Equation 4.74
- Ψ Constant used in Equation 4.32, and defined in Equation 4.33.
- Ω constant used in Equation 4.112, and defined in Equation 4.113
- ξ Constant used in the solution of differential equation during mud circulation

APPENDIX B

OVER-ALL HEAT TRANSFER COEFFICIENT FOR WELLS.

During the production of crudes, the fluid loses heat to the surrounding. Heat being transferred from the wellbore fluid to the earth has to overcome the resistances offered by the tubing wall, the tubing insulation (if any), the tubing-casing annulus, the casing wall, and the cement. This configuration is shown in Figure 21. The resistances are in series, and except for the annulus, the mechanism involved is conductive heat transfer. However, because of the presence of gas or liquid, convective heat transfer dominates in the annulus. In a wellbore/formation system, heat transfer does not really attain steady state. However, heat transfer rate variation is very slow and we assume steady state for a given time period and calculate the overall heat transfer coefficient based on that. In the next time period a new steady state is assumed and a new overall heat transfer coefficient is calculated.

At any steady state, the rate of heat flow through a wellbore per unit length of the well, ϕ , can be expressed as,

$$\phi = 2 \pi r_{\mu} U_{\mu} (T_{\mu} - T_{\mu}) \tag{A.1}$$

where U_{io} is defined as the over-all heat transfer coefficient based on the tubing outside surface area.





At steady state, heat flowing through each of the elements mentioned above will be same. This assumption of steady heat transfer allows us to write the rate of heat transfer across each element in terms of the temperature difference across the element and the resistance offered by that element. Thus, the rate of heat transfer between the flowing fluid and inside the tubing wall may be written as,

$$\phi = 2 \pi r_{\mu} h_{\ell} (T_{\ell} - T_{\mu})$$
(A.2)

Equation A.2 may be rewritten for the temperature drop across this element as,

$$T_f - T_u = \frac{\Phi}{2 \pi r_u h_t} \tag{A.3}$$

Noting that the sum of the temperature drops across all these elements is equal to the temperature difference between the fluid and the wellbore/earth interface, we can write,

$$T_{f} - T_{wb} = (T_{f} - T_{ii}) + (T_{ii} - T_{io}) + (T_{io} - T_{ins}) + (T_{ins} - T_{ci}) + (T_{ci} - T_{co}) + (T_{co} - T_{wb})$$
(A.4)

Or,

$$T_{f} - T_{wb} = \frac{\Phi}{2 \pi} \left[\frac{1}{r_{ii} h_{i}} + \frac{\ln(r_{io}/r_{ii})}{k_{i}} + \frac{\ln(r_{ins}/r_{io})}{k_{ins}} \right] + \left[\frac{1}{r_{ins}(h_{c} + h_{r})} + \frac{\ln(r_{co}/r_{ci})}{k_{cas}} + \frac{\ln(r_{wb}/r_{co})}{k_{cem}} \right]$$
(A.5)

$$T_{f} - T_{wb} = \frac{\Phi}{2 \pi r_{w} U_{to}}$$
 (A.6)

which is another form of Equation A.1, where U_{to} is given by,

$$\frac{1}{U_{io}} = \frac{r_{io}}{r_{ii}h_{i}} + \frac{r_{io}\ln(r_{io}/r_{ii})}{k_{i}} + \frac{r_{io}\ln(r_{ins}/r_{io})}{k_{ins}} + \frac{r_{io}}{r_{ins}(h_{c} + h_{r})} + \frac{r_{io}\ln(r_{co}/r_{ci})}{k_{cas}} + \frac{r_{io}\ln(r_{wb}/r_{co})}{k_{cem}}$$
(A.7)

Most of the terms in the above Equation can be easily computed except the fourth term. The fourth term, which represents the resistance to heat transfer offered by the annulus, is somewhat difficult to estimate. In most cases of petroleum production, the temperature difference across the annulus, is usually small and one need only to consider convective (natural) heat transfer. Unfortunately, no work on natural convection in vertical annular geometry is reported in the literature. This work adapts, as done by Hasan and Kabir (1991) and Willhite (1967), the correlation proposed by Dropkin and Sommerscales (1965) for heat transfer coefficient for natural convection in fluids between two vertical plates. Their correlation for h_c , expressed for our geometry, is

$$h_{c} = \frac{0.049 \ (Gr \ Pr)^{0.333} \ Pr^{0.074} \ k_{a}}{r_{ins} \ \ln\left(r_{ci}/r_{ins}\right)}$$
(A.8)

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Or,

The Grashof number Gr, in Equation A.8, is defined as,

$$Gr = \frac{(r_{ci} - r_{ins})^3 g \rho_a^2 \beta (T_{ins} - T_{ci})}{\mu_a^2}$$
(A.9)

The Grashof number reflects the extent of motion of the annulus fluid due to the natural convection. The density of the heated fluid next to the insulation layer is less than that next to the casing, creating Euoyancy force. The viscous force, working against the buoyancy, generates a circular motion of the fluid in the annulus. Prandtl number, Pr, is a measure of the interaction between the hydrodynamic boundary layer and the thermal boundary layer and is defined as,

$$Pr = \frac{c_{pa} \,\mu_a}{k_a} \tag{A.10}$$

Not all the components shown in Figure 21 is present in every wells. In addition, some of the elements may offer negligible resistance to heat flow. For example, in most oilwells, tubing insulation is absent. The high values of conductivity of metals, coupled with relatively thin tubing and casing walls, allows us to make the assumption that temperature drop across both the tubing and casing walls may be neglected. In that case, $T_{ui} = T_{to}$ and $T_{ci} = T_{co}$. With this assumption the expression for the overall heat transfer coefficient simplifies to,

$$U_{to} = \left[\frac{1}{h_c} + \frac{r_{to} \ln(r_{wb}/r_{co})}{k_{cem}}\right]^{-1}$$
(A.11)

We use this equation for estimating the overall heat transfer coefficient, U_{ω} . For most oilwells, it adequately represents the overall heat transfer coefficient. In some cases, it is possible to neglect annulus convection in determining the overall heat transfer coefficient. In that case, h_c should be replaced with the conductivity of annular fluid.

APPENDIX C

PROGRAM LISTING

PROGRAM WRITTEN BY MOHAMMAD MAHBUBUL AMEEN DEPARTMENT OF CHEMICAL ENGINEERING UNIVERSITY OF NORTH DAKOTA GRAND FORKS, ND

\$debug

program flow						
print *,'						
print *,'						
print *,' '						
print *,' '						
print *,' '						
print *,'	*****	****	******	******	*****	***
1*******						
print *,'	*****	******				
1*******						
print *,'	*					
1 * '						
print *,'	*	Chemical	Engr. Dep	tt.		
1 * '			C 1			
print *.'	*	University o	f North Da	akota.		
1 * '						

	print *,' 1 * '	* Grand Forks, North Dakota 58202.				
	print *,' 1 * '	*				
	print *,' 1 * '	*				
	print *,' 1 * '	* Welcome to Two-Phase Flow.				
	print *,' 1 *	*				
	print *,' 1 * '	*				
	print *,' leen *'	* This program is written by M. Mahbub Am				
	print *,' 1 *	* and supervised by Prof. A.R. Hasan				
	print *,' 1 * '	*				
	print *,' 1*******	*************				
	print *,' 1*******	*************				
	print *,' print *,'					
	print *,'Enter 1 i	f you want to continue'				
	read $(*,*)$ iii if (iii and 1) and	- CO1				
	n (m.eq. 1) go	If (111 .eq. 1) go to 601				
	print *' '					
	print *.'					
	print *,' '					
	print *,' '					
	print *,' '					
	print *,'					
	print *,' '					
	print *,' '					
	print *,'	You have decided not to continue. Bye bye.'				
	go to 602					
601	continue					
	print *,' '					
	print *,' '					
	print *,' '					
	print *.' '					
	print *,' '					

print *,' ' print *,' print * print *,' print *, print *, print *,' print *. You have decided to continue with this ' print *,' program. So now you have two options. You can ' print *,' ru., either your own problem or a sample ' print *,' problem. The choice 's yours. If you want' print *,' to run your own problem, then enter 2. ' print *,' otherwise for sample problem enter 3.' print *.' print *,' ' read (*,*) jij if (jjj .eq. 2) go to 605 if (jjj .eq. 3) go to 606 print *,' You have selected a wrong option.' print *,' Please enter 2 or 3 now.' print *,' ' print *,' ' print *.' print *,' ' print *,' ' read (*,*) jjj if (jjj .eq. 2) go to 605 if (jjj .eq. 3) go to 606 print *.' This is not a place for fun. Bye.'

go to 602 605 print *,' ' print *, print *,' Which condition is known to you ?' print *,' If you know bottom condition, enter 4' print *,' If you know surface condition, enter 5' print *,' print *.' print *,' print *.' ' print *,' ' read (*,*) kkk if (kkk .eq. 4) go to 607 if (kkk .eq. 5) go to 608 print *,' You have entered a wrong option. ' print *,' Please enter 4 or 5 now. print *,' . print *,' print *,' ' print *,' print *,' print *.' print *,' print *,' print *,' print *,' print *,' ' read (*,*) kkk if (kkk .eq. 4) go to 607 if (kkk .eq. 5) go to 608 print *,' This is not a place for fun. Bye.' go to 602 607 a=1.0 go to 610 608 a=(-1.0) 610 continue print *,' ' print *,' ' print *.' '

print *.' ' print *,' ' print *,' Now you will have to enter some' values. Please remember that this is' print *,' print *,' not a user friendly program, so if you' make any mistake, then you will have' print *,' print *.' to restart.' print *,' ' print *,'Enter the known pressure (in psia)' read (*,*) p print *,'Enter the known temperature (in ferenhite)' read (*,*) tt print *,'Enter the gor (in scf/stb)' read (*,*) gor print *,' Enter the diameter of the pipe (in ft)' print *,'[If annuli, then enter dia of outside pipe]' read (*,*) d print *,' Enter the diameter of inside pipe (in ft)' print *,' [If single pipe, then enter 0 (zero)]' read (*,*) din print *,'Enter the production rate (in stb/day)' read (*,*) prod print *,'Enter specefic gravity of gas' read (*,*) sgg print *,'Enter the specefic gravity of oil' read (*,*) sgl print *,'Enter the value of degree api' read(*,*) api print *,'Enter the roughness parameter epsilon' read (*,*) eps print *,'Enter the value of delta p (in +ve psia)' read (*.*) dp print *,'Enter depth of the well in feet ' read (*,*) depth print *,'Enter the value of dt/dz (in degree f/ foot, +ve) ' read (*,*) dtdz print *,'Enter the inclination angle with horizontal' read (*,*) alpha

print *,'Enter the value of surface tension (-----)' read (*,*) surten print *,' What is the direction of flow ?' print *,' If upward, enter 1 ' print *,' If downward, enter -1' read (*,*) cc print *,'If the flow is cocurrent, enter 1' print *,'But if it is countercurrent, enter -1' read (*,*) cc2 go to 666 print *,' ' print *,' ' print *,' print *,' 606 print *.' You have decided to go for the' print *.' sample problem. Now if would you like to' print *,' see bottom to surface calculation,' print *,' then enter 6, otherwise if you wish to' print *.' see surface to bottom calculation,' print *,' then enter 7.' print *,' ' print *,' ' print *.' ' print *,' ' print *,' ' print *.' ' print *.' ' print *,' ' print *.' ' print *.' ' print *,' ' read (*,*) 111 if (111 .eq. 6) go to 611 if (lll .eq. 7) go to 612 print *,' You have entered a wrong option.' print *.' Please enter 6 or 7 now.' print *.' '

print *,' ' print *,' ' print *,' ' print *,' ' print *,' ' print *,' ' print *,' ' print *,' ' read (*,*) lll if (lll .eq. 6) go to 611 if (lll .eq. 7) go to 612 print *,' This is not a place for fun. Bye.' go to 602

611 open (unit=51,file='bot.dat',status='old') read (51,*) p,tt,gor,d,din,prod,sgg,sgl,api,eps,dp,depth,dtdz,a,al 1pha,surten,cc,cc2 go to 666

612 open (unit=52,file='sur.dat',status='old') read (52,*) p,tt,gor,d,din,prod,sgg,sgl,api,eps,dp,depth,dtdz,a,al lpha,surten,cc,cc2

666 open (unit=8,file='res3.dat',status='old') write (8,*)' input data' write (8,552) *****************//) 552 format (' write (8,550) 550 format (3x,' pressure',4x,'temperature ',3x,'gas oil ratio',3x, 1' diameter',4x,'production',/) write (8,551)p,tt,gor,d,prod 551 format (5f15.6,/) write (8,553) 553 format (3x,' s.G.(Gas)',4x,'s.G.(Liquid)',3x,' alpha ',3x, 1' epsilon',4x,' delta p' \checkmark) write (8,554)sgg,sgl,alpha,eps,dp 554 format (5f15.6) write (8,555) 555 format (19x,' depth ',3x,' dt/dz '/) write (8.556)depth,dtdz

556 format (15x,2f15.6,///) write (8,*)' output data' write (8.557) 557 format (' ***************////)

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```
open (unit=6,file='res1.dat',status='old')
    write (6,500)
500 format (//,5x,'pressure',5x,' eg ',5x,'pipe length',5x,'para
     1meter nnn')
    write (6,504)
1====='/)
   g = 32.2
   gc=32.174
   degree=0.0174532*alpha
   dz=0.0
   wp=p
   wtt=tt
   sumh=0.0
   sumf=0.0
   sumv=0.0
   write (6,505) p,tt,dz
505 format (5x,f8.2,5X,f10.4,5X,f10.4,/)
50 continue
   call overall (p,tt,d,gor,prod,sgg,sgl,api,eps,g,gc,dpdzt,nnn,a,dpd
   1zh,dpdzf,dpdzv,degree,eg,din,surten,cc,cc2,alpha,valphat,vmix,vsg)
   dzdp=1.0/dpdzt
   dzdp01=dzdp
   p01=p
   tt01=tt
   tt=tt+dtdz*dzdp01*dp
   p=p-dp*a
   dpdzh1=dpdzh
   dpdzf1=dpdzf
   dpdzv1=dpdzv
   call overall (p,tt,d,gor,prod,sgg,sgl,api,eps,g,gc,dpdzt,nnn,a,dpd
   1zh,dpdzf,dpdzv,degree,eg,din,surten,cc,cc2,alpha,valphat,vmix,vsg)
   dzdp=1.0/dpdzt
   dzdp02=dzdp
   tt=tt01+dtdz*dzdp01*dp*0.5
   P=p01-dp*0.5*a
   dpdzh2=dpdzh
```

```
dpdzf2=dpdzf
dpdzv2=dpdzv
```

call overall (p,tt,d,gor,prod,sgg,sgl,api,eps,g,gc,dpdzt,nnn,a,dpd lzh,dpdzf,dpdzv,degree,eg,din,surten,cc,cc2,alpha,valphat,vmix.vsg) dzdp=1.0/dpdzt dzdp03=dzdp dpdzh3=dpdzh dpdzf3=dpdzf dpdzv3=dpdzv

dzdpa=(dzdp01+dzdp02+4.0*dzdp03)/6.0 dpdzha=(dpdzh1+dpdzh2+dpdzh3*4.0)/6.0 dpdzfa=(dpdzf1+dpdzf2+dpdzf3*4.0)/6.0dpdzva=(dpdzv1+dpdzv2+dpdzv3*4.0)/6.0

p=p01-dp*a tt=tt01+dtdz*dzdpa*dp dzv=(-dzdpa)*dp*a dz=dz+dzv

pipel=dz/sin(degree)
write (6,501) p,eg,pipel,nnn,valphat,vmix,vsg
501 format (5x,f8.2,5X,f10.4,5X,f10.4,11X,i2,3f10.6)

sumh=sumh+dpdzha sumf=sumf+dpdzfa sumv=sumv+dpdzva checka=depth-dzv if (dz .gt. checka) go to 745 go to 50

745 checkb=depth-dz

coeff=checkb/dzv p04=p tt04=tt p=p-0.5*dp*coeff*a tt=tt+dtdz*dzdpa*0.5*dp*coeff call overall (p,tt,d,gor,prod,sgg,sgl,api,eps,g,gc,dpdzt,nnn,a,dpd lzh,dpdzf,dpdzv,degree.eg,din,surten,cc,cc2,alpha,valphat,vmix,vsg)

dpp=dpdzt*checkb p=p04+dpp tt=tt04-dtdz*dpp*a/dpdzt dz=dz+checkb sumh=sumh+dpdzh sumf=sumf+dpdzf sumv=sumv+dpdzv suma=(sumh+sumf+sumv) frach=sumh/suma fracf=sumf/suma fracv=sumv/suma delp=abs(wp-p) delph=delp*frach delpf=delp*fracf delpv=delp*fracv

write (6,502)

```
write (8,560)
```

560 format (7x,'pressure',3x,' temperature',' parameter nnn',8x,'delp 1(h)',7x,'delp (f)',/) write (8,561)p,tt,wnnn,delph,delpf

```
561 format (5f15.5,//)
```

print *,' ' print *,' print *,' ' print *,' data entered by the user ' print *,' print *,' ' print *,' diame pressure temperature delta p 1ter' print *,' С 1---' С print *,wp,wtt,dp,d print *,' ' s.g.(gas) s.g.(liquid) production print *,' go 1r'----print *,' С С 1--' print *,sgg,sgl,prod,gor print *,' '

print *,' epsilon dt/ alpha depth ldz' с print *,' --------------1---' с print *,alpha,depth,eps,dtdz print *,' print *,' ' print *,' A very brief look at the results' print *,' (Conditons at the other end)' print *,' Detailed results are in res1.dat, res2.dat, res3.dat 1files' ****** print *,' 1***** ' print *,' ' print *,' dp/ presssure temperature nnn 1dz' print *,' С ---------------1---' с wnnn= nnn print *,p,tt,wnnn,dpdzt print *,' ' print *,' delta p(t) delta p(h) delta p(f)delt 1a p(v)'print *.' с ----------1-----' С print *,delp,delph,delpf,delpv go to 629 602 continue print *,' ' print *,' ' print *.' ' print *,' ' print *.' ' print *,' ' print *.' ' 629 continue end

\$debug

subroutine overall (p,tt,d,gor,prod,sgg,sgl,api,eps,g,gc,dpdzt,nnn 1,a,dpdzh,dpdzf,dpdzv,degree,eg,din,surten,cc,alpha,valphat,vmix,vs 1g)

call property (p,tt,d,gor,prod,sgg,sgl,api,vsg,vsl,rhog,rhol,fg,rs 1.din)

- print *, 'temperature in ferenhite =',tt,' free gas =',fgС С
 - print *, 'density of gas =',rhog,' density of liquid =',rho
- print *, 'velocity of gas =',vsg,'velocity of liquid =',vsl С

```
call viscosit (tt,visg,visl,rs)
```

call pattern (vsg,vsl,vmix,rhog,rhol,rhoc,rhom,visg,visl,d,eg,nnn, lg,degree,din,surten,cc,alpha,valphat)

print *,' eg =',eg,rhol,rhog,rhom С

> call gradient (vsg,vmix,rhol,rhom,rhoc,visg,visl,eps,d,eg,p,dpdzt, 1g,gc,nnn,a,dpdzh,dpdzf,dpdzv,degree,din,cc)

return end

\$debug

subroutine property (p,tt,d,gor,prod,sgg,sgl,api,vsg,vsl,rhog,rhol 1,fg,rs,din)

```
t = tt + 460.0
if (api .Le. 10.0) go to 103
qmo=650.0-11.0*api+(1.9E-7)*(api**5.0)
fl=(p*sgg)/t
```

if (il .ge. 1.0) go to 101 yg=0.43*fl**1.2-0.12*fl**4.0 go to 102 101 yg=0.56*alog10(fl)+0.315

102 continue

```
rs=((379.3*350.0*sgl)/qmo)*(yg/(1.0-yg))
go to 111
```

103 continue

rs=sgg*((p/18.0)*((10**(0.0125*api))/(10**(0.00091*tt))))**1.20481 193

- 111 fg=gor-rs if (fg .gt. 0.0) go to 112 rs=gor fg=0.0
- 112 pc=692.0-30.0*sgg-8.0*sgg**2.0 pr=p/pc tc=162.0+328.0*sgg tr=t/tc

prtr=pr/(tr**2.0) C if (tr .lt. 1.1 .or. prtr .gt. 1.0) go to 106

```
cm=0.51*tr**(-4.133)
cn=0.038-0.026*tr**(0.5)
if (tr .gt. 2.4) go to 104
zzz=0.0003
go to 105
104 zzz=0.0007
105 z= 1.0-cm*pr+cn*pr**2.0+zzz*pr**3.0
C 106 continue
```

c Calculate by some other method in this space

area=(3.1416*(d**2.0-din**2.0)/4.0) bo=0.972+0.000147*((rs*(sgg/sgl)**0.5)+1.25*tt)**1.175 r=10.73 print *,'area =',area,p vsg=(fg*prod*t*14.69*z)/(24.0*3600.0*520.0*p*area) vsl=(prod*bo*6.49*10.0**(-5.0))/area rhog=(29.0*sgg*p)/(z*r*t)

rhol=(350.4*sgl+rs*sgg*0.0764)/(5.615*bo) open (unit=7,file='res2.dat',status='old') write (7,505)p,tt,fl,yg,q:no.rs,fg,z,rhog,rhol,vsg,vsl 505 format (2x,12f9.3) return end

\$debug

subroutine pattern (vsg,vsl,vmix,rhog,rhol,rhoc,rhom,visg,visl,d,e 1g,nnn,g,degree,din,surten,cc,alpha,valphat)

c check for bubbly flow

```
valpha=1.53*((g*surten*((rhol-rhog)/(rhol**2)))**0.25)
c0=1.2
if (cc .eq. -1) c0=1.12
vmix=vsl+vsg
eg=vsg/(c0*vmix+cc*valpha)
rhom=rhol*(1.0-eg)+rhog*eg
```

c first check

check1=(0.429*vsl+cc*0.357*valpha)*sin(degree) if (check1 .gt. vsg) go to 150

```
c second check
```

```
de=d-din
check2=4.68*(de**0.48)*((g*((rhol-rhog)/surten))**0.5)*((surten/rh
lol)**0.6)*((rhol/visl)**0.08)
check3=vmix**1.12
if (check3 .gt. check2 .and. eg .lt. 0.52) go to 150
go to 151
```

c bubbly flow

```
150 if (vsg .le. 0.000000001) go to 152
nnn=1
go to 200
152 nnn=0
go to 200
```

```
no bubbly flow
с
 151 continue
С
     check for slug flow
     if (check3 .lt. check2) go to 170
     go to 171
С
     slug flow
170 nnn=2
    go to 199
     no slug flow
С
171 continue
     check for churn flow
с
    check9=3.1*(surten*g*(rhol-rhog)/(rhog**2.0))**0.25
    if (check9 .gt. vsg) go to 173
    go to 181
173 nnn=3
    c1=1.15
    if (alpha .le. 70) c1=1.2
    if (cc .eq. -1) c1=1.12
    go to 197
199 c1=1.2
    if (cc .eq. -1) c1=1.12
197 continue
     Calculation of v(alpha-t) follows, then eg and rhom
С
    anf=(((de**3.0)*g*(rhol-rhog)*rhol)**0.5)/visl
     aneo=(g*(de**2)*(rhol-rhog))/surten
     if (anf .gt. 250.0) go to 155
     if (anf .gt. 18.0) go to 156
     am=25.0
     go to 159
```

```
155 am=10.0
```

```
go to 159
```

```
156 am=69.0*(anf**0.35)
```

159 continue

ca2=0.345*(1.0-exp((-0.01)*anf/0.345))*(1.0-exp((3.37-aneo)/am))

```
valphat=(0.345+0.1*(din/d))*((sin(degree))**0.5)*((1+cos(degree))*
1*1.2)*((g*d*(rhol-rhog)/rhol)**0.5)
```

C print *,'valphat =',valphat, valpha

egt=vsg/(c1*vmix+cc*valphat) if (vsg .le. 1.312333) go to 166

```
qlsl=0.1*((c0*vmix+cc*valpha)/vsg)
qltl=1.0-qlsl
eg=qltl*egt+0.1
go to 167
```

- 166 qlsl=0.0762*(c0*vmix+cc*valpha) qltl=1.0-qlsl eg=qltl*egt+0.0762*vsg
- 167 rhom=rhog*eg+rhol*(1.0-eg) go to 200
- c Annular flow (no check needed)
- 181 nnn=4

```
xx=(vsg*rhog)/(vsg*rhog+vsl*rhol)
x=((rhog/rhol)**0.5)*(((1.0-xx)/xx)**0.9)*((visl/visg)**0.1)
eg=(1.0+x**0.8)**(-0.378)
```

```
vcgs=(vsg*visg*(rhog/rhol)**0.5)/surten
if (vcgs .ge. 0.0004) go to 185
e=0.0055*((vcgs*10000.0)**2.86)
go to 186
```

- 185 e=0.857*(log10(vcgs*10000.0))-0.2
- 186 continue
- c Another way of calculating eg

```
rhoc=(vsg*rhog+e*vsl*rhol)/(vsg+e*vsl)
```

200 continue return end \$debug

subroutine viscosit (tt,visg,visl.rs)

c calculation of viscosity

c dead oil

dla=1.54*10.0**8.0 dlm=3.12 Visdl=dla/(tt**dlm)

c live oil

С

ela=0.6*exp((-0.0032)*rs)+0.4*exp((-0.00011)*rs) elb=0.3*exp((-0.0035)*rs)+0.7*exp((-0.00022)*rs) visl=(ela*visdl**elb)*0.0006719 visl=0.000672

C Program needed to evaluate gas viscosity & liq. surface tension

visg=0.00000672 return end

\$debug

subroutine gradient (vsg,vmix,rhol,rhom,rhoc,visg,visl,eps,d,eg,p, ldpdzt,g,gc,nnn,a,dpdzh,dpdzf,dpdzv,degree,din,cc)

c gradient for annular flow

de=d-din if (nnn .ne. 4) go to 201 reg=(0.975*de*vsg*rhoc)/visg

```
fc=0.0791*(1.0+75.0*(1.0-eg))/(reg**0.25)
dpdzx=(+1.0)*((g*rhoc)+(2.0*fc*cc*rhoc*vsg**2.0)/(d*sin(degree)))/
1(gc*144.0*(1.0-(rhoc*vsg**2.0)/(p*144*gc*sin(degree))))
dpdzt=dpdzx*(-a)
open (unit=11, file='res4.dat'.status='old')
```

```
write(11,1001)p,rhoc,reg fc,dpdzt
```

```
1001 format (2x,2f8.3,1X,1f12.3,1X,2f10.3)
go to 300
```

```
c Common calculation for bubbly, slug and churn
```

```
201 re=(de*vmix*rhol)/visl
fa=((eps/de)**1.1098)/2.8257+(7.149/re)**0.8981
```

C f=(1.0/(4.0*log((eps/d)/3.7065-(5.0452*log(fa))/re)))**2.0 if (re .gt. 100000) go to 202 f=0.0791/(re**0.25) go to 203 202 f=0.046/(re**(0.2)) 203 if (nnn .le. 1) go to 211

c gradient for slug and churn flow

```
 dpdzf = (((-2.0 * cc*a) * f*vmix**2.0 * rhol*(1.0-eg))/(gc*d))/sin(degree) 
 dpdzh = ((a*(-rhom)*g)/gc*sin(0.0174532*90)) 
 dpdzv=0.0
```

```
dpdzt=(dpdzf+dpdzh+dpdzv)/144.0
```

```
open (unit=12,file='res5.dat',status='old')
write (12,1002)p,rhom,re,f,dpdzt
1002 format (2x,f10.3,2X,f8.5,2X,f12.3,1X,2f10.3)
go to 300
```

```
c gradient for bubblyflow
```

```
211 dpdzf=(((-2.0*cc*a)*f*vmix**2.0*rhom)/(gc*de))/sin(degree)
dpdzh=(a*(-rhom)*g)/gc*sin(0.0174532*90)
dpdzv=0.0
dpdzt=(dpdzf+dpdzh+dpdzv)/144.0
open (unit=13,file='res6.dat',status='old')
write (13,1003)p,rhom,re,f,dpdzt
1003 format (2x,f10.3,2X,f8.5,2X,f12.3,1X,2f10.3)
```

```
300 continue
return
end
```

\$debug

с subroutine super

> real kann.kcem.kfor dimension dqr'z(110,110).consp(110,110) dimension time(110).timed(110),timedd(110,110),timerd(110,110) dimension tdd(1:0:110)

This program calculates the fluid temperature С using superposition С

data cpliq, cpgas / 0.947, 0.200 / data kann, kcem, kfor / 0.383, 4.021, 0.83 / data dia, dcin, dcout, dcem / 0.2376, 0.5375, 0.5833, 0.75 / data alpha, beta, geograd / 0.04, 0.00011, 0.005926 / data p, gor, api, sgg / 113, 68, 34.3, 1.04 /

- С istep= # of time steps.
- C kstep= # length of depth step.
- nflag= 1 for analytical solution. С C
 - 2 for linear variation of phi.
- C mflag= 4 includes convection

5 excludes convection

print *,' enter the value of istep, kstep, nflag & mflag' read (*,*) istep,kstep,nflag,mflag

zbh=5400 tforbi=108 tottim=158

С

с viscosity and rhol should come from another subroutine.

visl=3.8rhol=62.3

С wg and wl comes from main prog.

> wg=0.00 wl = 2.46

wtot=wg+wl qual=wg/wtot cpm=cpgas*qual+cpliq*(1-qual)

klstep=zbh/kstep kk=kistep+2 do 19 k=2,kk,1dqdz(1,k)=0.0dqdz(2,k)=0.0continue

```
19
```

cons=-0.002978+1.006E-6*p+1.906E-4*wtot-1.047E-6*gor+3.229E-5*api+ 14.009E-3*sgg-0.3551*geograd

timed(1)=0.0timed(2)=0.0

delti=tottim/istep ii=istep+2 do 29 l=3,ii,1 time(l)=time(l-1)+delti

29 continue

```
do 401 l=3,ii,1
timed(l)=4*alpha*time(l)/(dcem**2)
```

```
do 402 m=1,1-1,1
timedd(l,m)=timed(l)-timed(m)
timerd(l,m)=timedd(l,m)**0.5
tdd(1,m) = (0.4063 + 0.5 * \log(timedd(1,m))) * (1+0.6/timedd(1,m))
if (timedd(1,m) .le. 1.5) Tdd(1,m)=1.12812*timerd(1,m)*(1-0.3*time
1rd(l,m)
```

```
402 continue
```

```
401 continue
```

do 777 i=3.ii,1

dz=0.0 tfluid=tforbi

do 888 j=2,kk,1

z=zbh-dz dt=5.00

100 Iter=0

go to 102 101 dt=dto+(dt-dto)*0.2102 continue kfor=0.83C if (z.lt. 1001) kfor= 1.4-0.00057*z if (mflag .eq. 5) go to 105 prnum=(cpliq*visl)/kann gras=((((3600**2)*32.2)/8)*((dcin-dia)**3)*(rhol**2)*beta*dt)/visl 1**2 convec=0.049*((prnum*gras)**0.333)*(prnum**0.074)*kann if (convec .le. kann) go to 106 go to 107 105 kfor=kfor*1.687 106 convec=kann 107 resann=((0.5*dia)*log(dcin/dia))/convec rescem=((0.5*dia)*log(dcem/dcout))/kcem uto=1/(rescem+resann) tfor=tforbi+geograd*(-dz) print *, cpm,wtot,kfor,tdd(i,i-1),dia,uto с afact=((cpm*wtot*3600)*(kfor+tdd(i,i-1)*(dia/2)*uto))/(3.1416*dia* 1kfor*uto) bfact=(kfor+(dia/2)*uto*tdd(i,i-1))/((dia/2)*uto)print *, afact, bfact С sum=0.0do 403 l=3,i,1 sum=sum+(dqdz(l-1,j)-dqdz(l-2,j))*tdd(i,l-2)403 continue if (nflag .eq. 2) go to 39

if (z .eq. zbh) go to 37 $dqdz(i,j)=((cpm^{*}(tfor-tfluid))/afact)+(tdd(i,i-1)^{*}dqdz(i-1,j)/bfac$
1t)-(sum/bfact)

go to 36

37 dqdz(i,j)=0.0

```
36 dto=dt
```

```
consp(i,j)=cons+((tdd(i,i-1)*dqdz(i-1,j))/(cpm*bfact))-(sum/(bfact 1*cpm))
```

```
tfact=afact*(-1.0/(778*cpm)+consp(i,j)+geograd)
expo=exp((-zbh+z)/afact)
if (expo .eq. 1.0) expo=0.99999
tfluid=tfor+tfact-expo*(tfact)
```

q1 = -dqdz(i,j)*wtot*3600

if (mflag .eq. 5) Go to 45

dt=(-dqdz(i,j)*wtot*3600)/((3.1416*dia)*convec)

go to 47

39 if (z .eq. zbh) go to 301 dqdz(i,j)=((cpm*(tfor-tfluid))/(afact*(zbh-z)))+(tdd(i,i-1)*dqdz(i 1-1,j)/bfact)-(sum/bfact) go to 302

```
301 \, dqdz(i,j)=0.0
```

```
302 dto=dt
```

```
ft=geograd-(tdd(i,i-1)*afact*dqdz(i-1,j))/(bfact*cpm)+(afact*sum)/
1(cpm*bfact)
```

```
tfact=afact*(-1.0/(778*cpm)+cons+ft)
expo=exp((-zbh+z)/afact)
if (expo .eq. 1.0) expo=0.99999
tfluid=tforbi-(zbh-z)*ft+tfact-expo*(tfact)
q1=-dqdz(i,j)*(zbh-z)*wtot*3600
```

```
if (mflag .eq. 5) go to 45
dt=(-dqdz(i,j)*(zbh-z)*wtot*3600)/((3.1416*dia)*convec)
```

47 continue

print *,z,dqdz(i,j) if (dt .le. 0.0000001) dt=0.00001 cky001=abs((dt-dto)/dt) iter=iter+1

if (iter .gt. 199) go to 45 if (cky001 .ge. 0.001) go to 101

45 continue

С

if (i .eq. ii) go to 10

```
if (z .eq. 5400) go to 10
if (z .eq. 5300) go to 10
if (z.eq. 5200) go to 10
if (z .eq. 5100) go to 10
if (z .eq. 5000) go to 10
if (z .eq. 4500) go to 10
if (z .eq. 4000) go to 10
if (z .eq. 3500) go to 10
if (z .eq. 3000) go to 10
if (z .eq. 2500) go to 10
if (z .eq. 2000) go to 10
if (z .eq. 1500) go to 10
if (z .eq. 1000) go to 10
if (z .eq. 500) go to 10
if (z .eq. 100) go to 10
if (z .eq. 000) go to 10
```

go to 12

- 10 open (unit=7,file='ress.dat',status='old') write(7,11)z,tfor,tfluid,dt,q1,afact,bfact,uto
- 11 format (8f12.5)
- 12 tfluid=tfluid-geograd*0.5*kstep
- 888 dz=dz+kstep

print *,' I =',i,' j =',j

777 dtime=dtime+delti

```
stop
end
```

\$debug

program dan

- c This program calculates the fluid temperature (in the tube &
- c in the annuli) using superposition. Numerical solution
- c wt & wa same.

real kfor dimension dqdz(220,110) dimension time(110),timed(110),timedd(110,110),timerd(110,110) dimension tdd(220,110),tft(500),tfa(500)

- c This program calculates the fluid temperaturec using superposition
- c Down the annuli & up the tube, mmm=+1
 c Down the tube & up the annuli, mmm=-1

data cpliq, cpgas / 0.400, 0.200 / data kfor, mmm / 1.30, -1 / data diat, diaa / 0.46875, 0.697916 / data alpha, geograd / 0.039, 0.0127 / data uta, utt / 32.1, 30.0 /

print *,' enter the value of istep and kstep ' read (*,*) istep,kstep

zbh=5000tforti=59.5 tai=75.0 tti=75.0 tottim=44.17 delz=50

Viscosity and rhol should come from another subroutine.

visl=3.8 rhol=62.3

С

C

wg and wl comes from main prog.

wg=0.00 wl=35.0

```
wtot=wg+wl
     qual=wg/wtot
     cpm=cpgas*qual+cpliq*(1-qual)
     klstep=zbh/kstep
     kk=klstep+2
     kkk=kk-2
     do 19 k=2.kk.1
     dqdz(1,k)=0.0
     dqdz(2,k)=0.0
19
     continue
     timed(1)=0.0
     timed(2)=0.0
    delti=tottim/istep
    ii=istep+2
    do 29 l=3,ii,1
    time(l)=time(l-1)+delti
    continue
 29
    do 401 l=3,ii,1
    timed(l)=4*alpha*time(l)/(diaa**2)
    do 402 m=1,1-1,1
    timedd(l,m)=timed(l)-timed(m)
    timerd(l,m)=timedd(l,m)**0.5
    tdd(1,m)=(0.4063+0.5*log(timedd(1,m)))*(1+0.6/timedd(1,m))
    if (timedd(l,m) .le. 1.5) tdd(l,m)=1.12812*timerd(l,m)*(1-0.3*time
    1rd(l,m)
402 continue
401 continue
    do 777 i=3,ii,1
    zbh=5000
    tforti=59.5
    tai=75.0
```

tti=75.0 tft(2)=tti

tfa(2)=77

z=0.0 dz=0.0

```
do 888 j=2,kkk,1
z=dz
z1=z+50
z2=z1+50
```

100 continue

```
tfor=tforti+geograd*dz
afact=((cpm*wtot*3600)*(kfor+tdd(i,i-1)*(diaa/2)*uta))/(3.1416*dia 1a*kfor*uta)
bfact=(kfor+(diaa/2)*uta*tdd(i,i-1))/((diaa/2)*uta)
bfact=(wtot*3600*cpm)/(2*3.1416*(diat/2)*utt)
```

```
sum=0.0
do 403 l=3,i,1
sum=sum+(dqdz(l-1,j)-dqdz(l-2,j))*tdd(i,l-2)
```

403 continue

С

С

```
dqdz(i,j)=((cpm*(tfor-tfa(j)))/afact)+(tdd(i,i-1)*dqdz(i-1,j)/bfac
1t)-(sum/bfact)
```

```
q1=-dqdz(i,j)*wtot*3600

con1=((afact*dqdz(i-1,j))/(bfact*cpm))*tdd(i,i-1)

con2=(afact/(cpm*bfact))*sum
```

```
if (mmm .Eq. -1) go to 22
```

```
tlam 1=(-1/(2*afact))+(1/(2*afact))*sqrt(1+4*((diaa/2)*uta*tdd(i,i-11)+kfor)*(((diat/2)*utt)/((diaa/2)*uta*kfor)))
```

```
tlam2=(-1/(2*afact))-(1/(2*afact))*sqrt(1+4*((diaa/2)*uta*tdd(i,i-11)+kfor)*(((diat/2)*utt)/((diaa/2)*uta*kfor)))
```

talpha=-((tai-tforti-con1+con2)*tlam2*exp(tlam2*zbh)+geograd*(1-tl 1am2*bbfact))/(tlam1*(exp(tlam1*zbh))*(1-tlam2*bbfact)-tlam2*(exp(t 1lam2*zbh))*(1-tlam1*bbfact))

```
tbeta= ((tai-tforti-con1+con2)*tlam1*exp(tlam1*zbh)+geograd*(1-tla
lm1*bbfact))/(tlam1*(exp(tlam1*zbh))*(1-tlam2*bbfact)-tlam2*(exp(tl
lam2*zbh))*(1-tlam1*bbfact))
```

print *,tlam1,tlam2,talpha,tbeta

if (j .eq. 2) then

tft(j)=tai

tft(j+1)=(1-tlam1*bbfact)*talpha*exp(tlam1*z1)+(1-tlam2*bbfact)1)*tbeta*exp(tlam2*z1)+geograd*z+tforti+con1-con2

 $tft(i+2) = -((delz^{*}2)/(afact^{*}bbfact))^{*}(bbfact^{*}(tft(i+1)-tft(i)))$ 1/delz-tft(j)+tforti+geograd*z+con1-con2)+2*tft(j+1)-tft(j)

der0 = (tft(j+1)-tft(j))/delztfa(j)=tft(j)+bbfact*der0tfa(j+1)=tft(j+1)+bbfact*der0

der1 = (tft(j+2)-tft(j+1))/delztfa(j+2)=tft(j+2)+bbfact*der1

else

 $tft(j+2) = -((delz^{*}2)/(afact^{*}bbfact))^{*}(bbfact^{*}(tft(j+1)-tft(j)))$ 1/delz-tft(j)+tforti+geograd*z+con1-con2)+2*tft(j+1)-tft(j)

```
der1 = (tft(j+2)-tft(j+1))/delz
   tfa(j+2)=tft(j+2)+bbfact*der1
   print *,z,der1
end if
```

tft=talpha*exp(tlam1*z1)+tbeta*exp(tlam2*z)+geograd*z1+bbfact*geog С С 1rad+tforti+con1-con2

go to 24

22 tlam1=(1/(2*afact))+(1/(2*afact))*sqrt(1+4*((diaa/2)*uta*tdd(i,i-1)))1)+kfor)*(((diat/2)*utt)/((diaa/2)*uta*kfor)))

tlam2=(1/(2*afact))-(1/(2*afact))*sqrt(1+4*((diaa/2)*uta*tdd(i,i-1)))1)+kfor)*(((diat/2)*utt)/((diaa/2)*uta*kfor)))

tgamma=-((tti-tforti+bbfact*geograd-con1+con2)*tlam2*exp(tlam2*zbh 1)+geograd)/(tlam1*(exp(tlam1*zbh))-tlam2*(exp(tlam2*zbh)))

tdelta= ((tti-tforti+bbfact*geograd-con1+con2)*tlam1*exp(tlam1*zbh 1)+geograd)/(tlam1*(exp(tlam1*zbh))-tlam2*(exp(tlam2*zbh)))

С tft=tgamma*exp(tlam1*z)+tdelta*exp(tlam2*z)+geograd*z-bbfact*geogr

c 1ad+tforti+con1-con2

c tfa=(1+tlam1*bbfact)*tgamma*exp(tlam1*z)+(1+tlam2*bbfact)*tdelta*e

c 1xp(tlam2*z)+geograd*z+tforti+con1-con2

if (j .eq. 2) then

tft(j)=tti

tft(j+1)=tgamma*exp(tlam1*z1)+tdelta*exp(tlam2*z1)+geograd*z1-1bbfact*geograd+tforti+con1-con2 !-0.0062

 $tft(j+2)=((delz^{*}2)/(afact^{*}bbfact))^{*}(bbfact^{*}(tft(j+1)-tft(j))/1delz+tft(j)-tforti-geograd^{*}z2-con1+con2)+2^{*}tft(j+1)-tft(j)$

print *,j,tft(j+1),tft(j+2)

der00=(tft(j+1)-tft(j))/delzder01=(tft(j+2)-tft(j+1))/delzder0=der00-(der01-der00)

print *,der00,der01,der0

tfa(j)=tft(j)+bbfact*der0tfa(j+1)=tft(j+1)+bbfact*der00tfa(j+2)=tft(j+2)+bbfact*der01

else

С

 $tft(j+2)=((delz^{*}2)/(afact^{bbfact}))^{*}(bbfact^{*}(tft(j+1)-tft(j))/1delz+tft(j)-tforti-geograd^{*}z2-con1+con2)+2^{*}tft(j+1)-tft(j)$

der 1 = (tft(j+2)-tft(j+1))/delztfa(j+2) = tft(j+2) + bbfact*der1

end if

24 open (unit=7,file='ress.dat',status='old')

if (j.eq. 2) then

write (7,13) z,tfa(j),tft(j),con1.con2

13 format (5f12.5)

140

else

```
write(7,11)z2,tfa(j+2),tft(j+2),con1,con2
```

```
11 format (5f12.5)
```

end if

888 dz=dz+kstep

print *,' I =',i,' j =',j

777 dtime=dtime+delti

stop end

\$debug

program da

- c This program calculates the fluid temperature (in the tube &
- c in the annuli) using superposition.

C wt & wa same.

```
teal kfor
dimension dqdz(100,60)
dimension time(100),timed(100),timedd(100,100),timerd(100,100)
dimension tdd(100,100)
```

```
c This program calculates the fluid temperature using superposition
```

```
c Down the annuli & up the tube, mmm=+1c Down the tube & up the annuli, mmm=-1
```

```
data cpliq, cpgas / 0.400, 0.200 /
data kfor / 1.30 /
data diat, diaa / 0.46875, 0.697916 /
data alpha, geograd / 0.039, 0.0127 /
data uta, utt / 32.1, 30.0 /
```

```
print *,' enter the value of istep, kstep and mmm' read (*,*) istep,kstep,mmm
```

```
zbh=15000
tforti=59.5
tai=75.0
```

tti=75.0 tottim=44.17

C viscosity and rhol should come from another subroutine.

visl=3.8 rhol=62.3

C wg and wl comes from main prog.

wg=0.00 wl=35.0

```
wtot=wg+wl
qual=wg/wtot
cpm=cpgas*qual+cpliq*(1-qual)
```

```
klstep=zbh/kstep
kk=klstep+2
do 19 k=2,kk,1
dqdz(1,k)=0.0
dqdz(2,k)=0.0
```

```
19 continue
```

timed(1)=0.0timed(2)=0.0

delti=tottim/istep ii=istep+2

do 29 l=3,ii,1 time(l)=time(l-1)+delti

29 continue

do 401 l=3,ii,1

timed(1)=4*alpha*time(1)/(diaa**2)

do 402 m=1,1-1,1

timedd(l,m)=timed(l)-timed(m)
timerd(l,m)=timedd(l,m)**0.5

tdd(1,m) = (0.4063 + 0.5*log(timedd(1,m)))*(1+0.6/timedd(1,m))

if (timedd(l,m) .le. 1.5) tdd(l,m)=1.12812*timerd(l,m)*(1-0.3*time lrd(l,m))

402 continue

401 continue

do 777 i=3,ii,1

```
zbh=15000
tforti=59.5
tai=75.0
tfa=tai
```

z=0.0 dz=0.0

do 888 j=2,kk,1

z=dz

100 continue

```
tfor=tforti+geograd*(dz)
```

```
afact=((cpm*wtot*3600)*(kfor+tdd(i,i-1)*(diaa/2)*uta))/(3.1416*dia 1a*kfor*uta)
bfact=(kfor+(diaa/2)*uta*tdd(i,i-1))/((diaa/2)*uta)
bbfact=(wtot*3600*cpm)/(2*3.1416*(diat/2)*utt)
```

sum=0.0do 403 l=3,i,1 sum=sum+(dqdz(l-1,j)-dqdz(l-2,j))*tdd(i,l-2)

403 continue

dqdz(i,j)=((cpm*(tfor-tfa))/afact)+(tdd(i,i-1)*dqdz(i-1,j)/bfact)-1(sum/bfact)

q1 = -dqdz(i,j) * wtot * 3600

con1=(((afact*dqdz(i-1,j))/(bfact*cpm)))*tdd(i,i-1)
con2=(afact/(cpm*bfact))*sum

if (mmm .eq. -1) go to 22

tlam l=(-1/(2*afact))+(1/(2*afact))*sqrt(1+4*((diaa/2)*uta*tdd(i,i-11)+kfor)*(((diat/2)*utt)/((diaa/2)*uta*kfor)))

tlam2=(-1/(2*afact))-(1/(2*afact))*sqrt(1+4*((diaa/2)*uta*tdd(i,i-11)+kfor)*(((diat/2)*utt)/((diaa/2)*uta*kfor)))

talpha=-((tai-tforti-con1+con2)*tlam2*exp(tlam2*zbh)+geograd*(1-tl 1am2*bbfact))/(tlam1*(exp(tlam1*zbh))*(1-tlam2*bbfact)-tlam2*(exp(t 1lam2*zbh))*(1-tlam1*bbfact))

tbeta= ((tai-tforti-con1+con2)*tlam1*exp(tlam1*zbh)+geograd*(1-tla 1m1*bbfact))/(tlam1*(exp(tlam1*zbh))*(1-tlam2*bbfact)-tlam2*(exp(tl 1am2*zbh))*(1-tlam1*bbfact))

tft=talpha*exp(tlam1*z)+tbeta*exp(tlam2*z)+geograd*z+bbf ct*geogra 1d+tforti+con1-con2

tfa=(1-tlam1*bbfact)*talpha*exp(tlam1*z)+(1-tlam2*bbfact)*tbeta*ex 1p(tlam2*z)+geograd*z+tforti+con1-con2

go to 24

22 tlam1=(1/(2*afact))+(1/(2*afact))*sqrt(1+4*((diaa/2)*uta*tdd(i,i-1)+kfor)*(((diat/2)*utt)/((diaa/2)*uta*kfor)))

tlam2=(1/(2*afact))-(1/(2*afact))*sqrt(1+4*((diaa/2)*uta*tdd(i,i-1)+kfor)*(((diat/2)*utt)/((diaa/2)*uta*kfor)))

tgamma=-((tti-tforti+bbfact*geograd-con1+con2)*tlam2*exp(tlam2*zbh 1)+geograd)/(tlam1*(exp(tlam1*zbh))-tlam2*(exp(tlam2*zbh)))

tdelta= ((tti-tforti+bbfact*geograd-con1+con2)*tlam1*exp(tlam1*zbh 1)+geograd)/(tlam1*(exp(tlam1*zbh))-tlam2*(exp(tlam2*zbh)))

tft1=tft

tft=tgamma*exp(tlam1*z)+tdelta*exp(tlam2*z)+geograd*z-bbfact*geogr lad+tforti+con1-con2

tfa=(1+tlam1*bbfact)*tgamma*exp(tlam1*z)+(1+tlam2*bbfact)*tdelta*e 1xp(tlam2*z)+geograd*z+tforti+con1-con2

c 23 print *,afact,tlam1,tlam2,tgamma,tdelta

- 24 open (unit=7,file='ress.dat',status='old') write(7,11)z,tfor,tft,tfa,dqdz(i,j).con1.con2
- 11 format (7f12.5)

888 dz=dz+kstep

print *,' I =',i,' j =',j

777 dtime=dtime+delti

stop

end

APPENDIX D

PROFILES IN A 8000 FT OILWELL

TABLE 5Wellbore and Fluid Data During Production	
Well Depth , ft	8000
Production Rate, lb/hr	8856
Tube Diameter, in	2.875
Casing Diameter, in	7.0
Wellbore Diameter, in	9.0
Specific Gravity of Crude, API	34.3
Formation Thermal Conductivity, Btu/hr ft F	0.83
Cement Thermal Conductivity, Btu/hr ft F	4.021
Annular Thermal Conductivity, Btu/hr ft F	0.383
Specific Heat of fluid, Btu/lb F	0.947
Surface Earth Temperature, F	76
Geothermal Gradient,	0.005926



Temperature Profile in the Wellbore Constant Psi, With Convection

Figure 22. Temperature vs. Depth in a 8000 ft Wellbore (Constant ψ and With Convection)



Figure 23. Heat Flux vs. Depth in a 8000 ft Wellbore (Constant ψ and With Convection)

Temperature Profile in the Wellbore Linear Phi, With Convection



Figure 24. Temperature vs. Depth in a 8000 ft Wellbore (Linear Heat Flux and With Convection)



Figure 25. Heat Flux vs. Depth in a 8000 ft Wellbore (Linear Heat Flux and With Convection)



Figure 26. Temperature vs. Depth in a 8000 ft Wellbore (Constant ψ and Without Convection)



Figure 27. Heat Flux vs. Depth in a 8000 ft Wellbore (Constant ψ and Without Convection)



Figure 28. Temperature vs. Depth in a 8000 ft Wellbore (Linear Heat Flux and Without Convection)

Heat Flux Profile in the Wellbore Linear Phi, Without Convection



Figure 29. Heat Flux vs. Depth in a 8000 ft Wellbore (Linear Heat Flux and Without Convection)

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