CHARACTERIZATION OF GAS-LIQUID-LIQUID TUBULAR REACTORS

BY

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To

Georgios and Stavroula

who took considerable pain

to raise and educate me.
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ABSTRACT

In this investigation, the impact of gas flux and physical properties of dispersed phase constituents on unsteady and steady-state drop size distributions of dispersed phase drops in gas-liquid-liquid tubular reactors is quantified. Experiments were performed with three model ternary dispersed phase systems including linseed oil+trichloroethylene-water-air, cottonseed oil+trichloroethylene-water-air and dibenzyl ether-water-air. The relative importance of dispersed phase viscosity, interfacial tension and the density difference between the two liquid phases are assessed independently. The physical properties of the dispersed phase are shown to have a significant impact on the steady-state mean drop size and drop size distribution at low gas fluxes. However, at high gas fluxes, the effect of dispersed phase physical properties is less significant. A new theory is proposed to explain the functional dependence of the maximum and minimum drop sizes, stable against break-up and coalescence, on the energy dissipation rate. Gas flux and physical properties of the dispersed phase constituents were not found to have a significant effect on the rate at which steady-state is reached. The existence of a similarity variable which is very helpful in estimating the drop size distribution as a function of time is identified. Mean drop size and drop size distributions are correlated using theoretical models for drop break-up and coalescence in turbulent flow fields and these results are compared with stirred tank analogues.
TABLE OF CONTENTS

Acknowledgements........................................................................................................... i
Abstract............................................................................................................................. ii
Table of Contents............................................................................................................... iii
List of Tables...................................................................................................................... v
List of Figures................................................................................................................... vi

1.0. INTRODUCTION........................................................................................................... 1

PART I - Theoretical
2.0. LITERATURE REVIEW............................................................................................... 6
  2.1. Bubble Formation...................................................................................................... 7
  2.2. Terminal Velocities of Bubbles............................................................................... 11
  2.3. Gas-Liquid Systems............................................................................................... 14
  2.4. Break-up and Coalescence of Drops in Turbulent Flow Fields-The Effect of Energy Dissipation......................................................................................... 17
  2.5. Dispersion Phenomena in Agitated Vessels......................................................... 20
  2.6. Objectives............................................................................................................... 23

PART II - Experimental
3.0. EXPERIMENTAL....................................................................................................... 26
  3.1. Description of the Experimental Apparatus......................................................... 26
  3.2. Experimental Procedure....................................................................................... 28

PART III - Results and Discussion
4.0. RESULTS.................................................................................................................. 37
  4.1. Unsteady State Drop Size Distributions............................................................... 37
  4.2. Steady State Drop Size Distributions................................................................... 38
5.0. DISCUSSION............................................................................................................. 69
  5.1. Gas Holdup............................................................................................................ 69
  5.2. Drop Size Distribution-Steady State................................................................... 71
  5.3. Drop Size Distribution-Unsteady State................................................................ 72
    5.3.1. Dynamics of Drop Breakage in Tubular Reactors........................................... 73
    5.3.2. Similarity and Probability of Drop Breakage.................................................. 75
    5.3.3. Comparison with Agitated Vessels (Unsteady State)...................................... 78
  5.4. Maximum, Minimum and Mean Drop Sizes....................................................... 79
    5.4.1. Superficial Gas Velocity.................................................................................. 79
    5.4.1.1. The Prediction of Mean Drop Size in Tubular Reactors for Gas-Liquid-Liquid Systems........................................................................................................... 79
    5.4.1.2. Experimental Results.................................................................................. 83
    5.4.2. Viscosity........................................................................................................... 90
    5.4.3. Interfacial Tension........................................................................................... 92
    5.4.4. Dispersed Phase Concentration...................................................................... 94
5.4.5. Density................................................................. 95
5.4.6. Other Parameters.................................................. 96
5.5. Comparison with Agitated Vessels (Steady State)............ 97

6.0. CONCLUSIONS........................................................... 100

7.0. RECOMMENDATIONS.................................................... 103

8.0. NOTATION............................................................... 105

9.0. REFERENCES........................................................... 108

APPENDICES

APPENDIX A
Calculation of the Energy Dissipation in Tubular Reactors........... 116

APPENDIX B
Maximum Drop Size stable against Break-up at Low Gas Fluxes........ 118

APPENDIX C
Minimum Drop Size stable against Coalescence at Low Gas Fluxes.... 121

APPENDIX D
Relationship Between Superficial Gas Velocity and Terminal
Velocity of Bubbles at Low Gas Fluxes.................................... 124
# LIST OF TABLES

<table>
<thead>
<tr>
<th>Table</th>
<th>Description</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Physical properties of gas-liquid-liquid systems</td>
<td>33</td>
</tr>
<tr>
<td>2</td>
<td>Sources and purities of chemicals</td>
<td>34</td>
</tr>
<tr>
<td>3</td>
<td>Repeatability of the procedure</td>
<td>35</td>
</tr>
<tr>
<td>4</td>
<td>Steady state maximum, minimum and Sauter mean diameters</td>
<td>40</td>
</tr>
<tr>
<td>5</td>
<td>Slopes of various systems in both flow regimes</td>
<td>41</td>
</tr>
<tr>
<td>6</td>
<td>The effect of relative density</td>
<td>42</td>
</tr>
</tbody>
</table>
LIST OF FIGURES

Figure 1: Schematic diagram of experimental apparatus................. 30
Figure 2: Detailed sketch of bubble column and gas chamber......... 31
Figure 3: Calibration curves for the rotameters........................... 32
Figure 4: Gas holdup in three phase tubular reactors.................... 43
Figure 5: Sample size distributions for dispersed phase drops....... 44
Figure 6: Sample cumulative drop size distributions...................... 45
Figure 7: Unsteady state cumulative volume distributions for the dibenzyl ether system ($u_g=0.1$ cm/sec)...................... 46
Figure 8: Unsteady state cumulative volume distributions for the cottonseed oil system ($u_g=0.4$ cm/sec).......................... 47
Figure 9: Unsteady state cumulative volume distributions for the linseed oil system ($u_g=2$ cm/sec)............................. 48
Figure 10: The effect of superficial gas velocity, $u_g$, on the unsteady state mean drop size (linseed oil system).............. 49
Figure 11: The effect of interfacial tension, $\sigma$, on the unsteady state mean drop size.................................................. 50
Figure 12: The effect of relative viscosity, $\mu_d/\mu_c$, on the unsteady state mean drop size................................................. 51
Figure 13: The functional dependence of the transition probability function $\Gamma(v)$ on drop size (dibenzyl ether system, $u_g=0.1$ cm/sec)................................................................. 52
Figure 14: The functional dependence of the transition probability function $\Gamma(v)$ on drop size (linseed oil system, $u_g=2$ cm/sec)........................................................................... 53
Figure 15: The similarity of the unsteady state cumulative volume size distributions (dibenzyl ether system, $u_g=0.1$ cm/sec)................................................................. 54
Figure 16: The similarity of the unsteady state cumulative volume size distributions (linseed oil system, \( u_g = 2 \text{ cm/sec} \))

Figure 17: A comparison between the rate at which steady state drop size distributions are approached in tubular reactors and agitated vessels

Figure 18: The effect of superficial gas velocity on the steady state Sauter mean diameter of dispersed liquid drops

Figure 19: Parity plot for Equation 5.16

Figure 20: The dependence of maximum and minimum drop sizes on the energy dissipation rate (cottonseed oil system)

Figure 21: The dependence of maximum and minimum drop sizes on the energy dissipation rate (linseed oil system)

Figure 22: The effect of energy dissipation rate on the breadth of drop size distributions for the linseed oil system

Figure 23: The effect of energy dissipation rate on the breadth of drop size distributions for the cottonseed system

Figure 24: The effect of energy dissipation rate on the size spectrum of dispersed phase drops (cottonseed oil system)

Figure 25: The effect of energy dissipation rate on the cumulative size spectrum of dispersed phase drops (linseed oil system)

Figure 26: The effect of relative viscosity, \( \mu_d / \mu_c \), on the steady state Sauter mean drop diameter

Figure 27: The effect of volume fraction, \( \phi \), of dispersed phase on the steady state Sauter mean diameter (linseed oil system)

Figure 28: The effect of volume fraction, \( \phi \), of the dispersed phase on the steady state Sauter mean diameter (cottonseed oil system)
Figure 29: A comparison of the performance of tubular reactors (experimental data) and agitated vessels (Equations 2.13 and 2.14).
1.0 INTRODUCTION

The dispersion of one phase in another phase in multiphase systems is of great importance in chemical engineering, specifically for processes involving heat and mass transfer. Moreover, hydrodynamic effects arising in these systems play an important role in determining the kinetic schemes and rates of transport processes associated with these systems. The design of multiphase reactors requires the study of both hydrodynamic effects and phase interactions. Studies related to two-phase systems are numerous. However, phase interactions associated with gas-liquid-liquid systems have received comparatively little attention even though these systems arise frequently in chemical engineering applications. For example, the reaction environment and kinetic schemes associated with the direct hydrogenation of bitumen and coal liquids are complex. Apart from vapour and solid phases, two liquid phases may also co-exist. Recently, Shaw et.al. (1988) carried out experiments using a model coal liquefaction solvent (pyrene-tetratin) and showed that two liquid phases can arise under coal liquefaction conditions. The transfer of hydrogen from the gas phase to the dispersed liquid phase has been identified as a potential rate limiting phenomenon in coal liquefaction. As the mass transfer rate limitation occurs at the liquid-liquid interface, mean drop size distribution becomes a key variable. In addition, the hydrogen bubbles may impose turbulent flow conditions while passing through the reactor. The criteria for bubble/drop
stability in isotropically turbulent two phase flow have been defined statistically by Thomas (1981). Such conditions are sufficiently intense to cause drop splitting and the generation of large interfacial areas, depending on the availability of hydrogen.

Phase interactions arising in these systems have been investigated in the present work using various physical models. The impact of gas-bubble liquid-drop interaction on the steady-state liquid-liquid interfacial area is emphasized. Moreover, the simultaneity of drop breakage and transport processes (heat and mass transfer) brings about the need for knowledge of drop size and drop size distribution as a function of time.

Due to the complexity of gas-liquid-liquid systems several characteristics related to these systems have been assessed and discussed independently. Bubble generation and motion are among the most important parameters influencing the operation of the phase-contacting equipment. As the bubbles move upwards, turbulent conditions are imposed throughout the reactor. Bubbles with different sizes have different terminal velocities and impose turbulent conditions with various levels of intensity. The rate of the energy dissipation is related to the intensity of turbulence and controls drop splitting. It also has a severe impact on the generation of interfacial area. Physical properties of the liquids are also expected to have a significant impact on the steady-state drop size and drop size distribution. The impact of various physical properties including
interfacial tension, viscosity and density is discussed. The behavior of gas-liquid-liquid systems in tubular reactors may be compared with liquid-liquid systems in agitated vessels by considering the respective roles of the gas-phase and the impeller in these reactor types. Thus, characteristics related to liquid-liquid systems in agitated vessels may provide some understanding of the behavior of gas-liquid-liquid systems in tubular reactors.

The various characteristics mentioned above are summarized in a generalized correlation for evaluating the steady state interfacial area in gas-liquid-liquid tubular reactors. It should be noted that the proposed correlation includes a broad range of operating conditions and dispersed phase properties. Also an attempt is made to model transient drop break-up so that the transient interfacial area can be known as a function of time. A new theory is proposed which is intended to elucidate the phase interactions related to gas-liquid-liquid systems in tubular reactors.

These various aspects may be used in establishing criteria for the design of G-L-L tubular reactors. It also improves the reactor operation and process control strategies. Once design criteria have been established, the following improvements over the existing tubular reactors are expected:

- a narrower residence time distribution for the dispersed phase
- improved gas-liquid mass transfer rates
- a reduced gas flux requirement
So far, the mechanism of drop splitting and the implications for the design and operation of industrial gas-liquid-liquid tubular reactors are unresolved even if they arise in chemical engineering applications. The present work sheds light on phase interactions arising in gas-liquid-liquid tubular reactors and proposes the further investigation which has to be done until these reactors might be applied efficiently in industry.
PART I

THEORETICAL
2.0 LITERATURE REVIEW

Phase interactions in gas-liquid, gas-solid, liquid-liquid and gas-liquid-solid systems have been the subjects of numerous experimental and theoretical investigations (Kumar et al., 1976; Kito et al., 1976; Begovich and Watson, 1978; Hikita et al., 1980; Kato et al., 1982; Stamatoudis and Tavlarides, 1985). However, phase interactions arising in gas-liquid-liquid systems have received comparatively little attention, even though such three phase systems can arise in hydrorefining processes (Shaw et al., 1988), pyrometallurgical processes (Guthrie and Bradshaw, 1969) and electroorganic synthesis. Yoshida and Yamada (1972) first measured the average diameter of kerosene drops dispersed in bubble columns which were operated in batch mode with respect to both continuous and dispersed phase liquids. The Sauter mean diameter was correlated as a function of power consumption per unit mass of liquid and volume fraction of dispersed phase liquid. The mean drop size was found to be roughly proportional to the power input (power input$^{-0.5}$) and to the volume fraction of the dispersed liquid phase (volume fraction$^{0.25}$). They also found that tubular reactors are more efficient contactors than agitated reactors for liquid-liquid systems as far as mean drop size is concerned. However, they did not consider the effect of physical properties of the dispersed and continuous phases. An empirical correlation relating mean drop size to column diameter and dispersed phase viscosity was
subsequently proposed by Hatate (1976).

Recently, Kato et al. (1984) carried out experiments in a multistage bubble column to simulate gas-liquid-liquid systems. The Sauter mean drop diameter was correlated as a function of superficial gas velocity, liquid velocity and free area of horizontal baffle plates. The operation of their column was continuous with respect to all three phases. However, it was unclear whether or not steady state was reached as far as mean drop size is concerned.

The behavior of gas-liquid-liquid systems is poorly characterized in the open literature. To gain a better understanding of the behavior of G-L-L systems, the behavior of gas-liquid systems in tubular reactors and liquid-liquid systems in agitated vessels are reviewed. In addition, bubble formation at orifices and drop break-up in turbulent flow fields may shed light on the controlling mechanisms of drop break-up in G-L-L systems.

### 2.1. Bubble Formation

Bubble formation is one of great importance in the operation of phase-contacting equipment for gas-liquid, gas-liquid-liquid and gas-liquid-solid systems. A variety of atomizers, spray nozzles and orifices have been devised for bubble generation. Each of these devices has different characteristics which have been reviewed (Kumar and Kuloor, 1970; Lane and Green, 1956; Clift et al., 1978). The various bubble sizes
generated by the above devices play an important role in determining the kinetic schemes and rates of transport processes associated with multi-phase systems (Shaw, 1985). An examination of the various parameters influencing bubble generation will shed light in understanding the operation of such multi-phase systems.

In the present work, bubbles are formed at a multi-nozzle plate located at the bottom of the column. When a bubble is formed at such a nozzle, the pressure within the bubble decreases due to its upward motion. Thus the gas rate may vary with time (Clift et al., 1978). Therefore, we can consider two distinct cases for bubble generation. When there is a relatively high pressure drop between the gas reservoir and the nozzle, the pressure fluctuations due to bubble formation are insignificant. In this case the gas flow can be taken constant and the bubbles are formed under constant flow conditions. The other extreme situation is taken when the chamber volume is very large by comparison with the bubbles being formed. This case, where the bubble generation does not change the pressure in the chamber, corresponds to bubble formation under constant pressure conditions. The intermediate range is called bubble generation under intermediate conditions. The division of bubble formation into three different mechanisms is quite fundamental since it allows for a broad range of bubble volumes at the same gas flow rate. Bubble volumes formed under constant pressure conditions are several times larger than those under
constant flow conditions at the same flow rate (Clift et al., 1978).

Davidson and Schuler (1960) were the first to distinguish these three bubble formation regimes. Tsuge and Hibino (1983) reviewed various correlations of bubble volume and classified the phenomenon of bubble generation into three groups using the dimensionless capacitance group $N_c$ proposed by Hughes et al. (1955).

1. $N_c < 1$  
   Constant flow conditions

2. $1 < N_c < 9$  
   Intermediate conditions

3. $N_c > 9$  
   Constant pressure conditions

where,

$$N_c = \frac{4V_c g \rho_l}{\pi D_o^2 P_h}$$

and $V_c$ is the volume of the chamber, $g$ the gravitational acceleration constant, $\rho_l$ the density of the liquid, $D_o$ the diameter of the orifice and $P_h$ the hydrostatic pressure at the orifice plate.

Davidson and Schuler (1960) also divided each of the above ranges into three subranges:

1. $N_{w} < 2.4(N_c-1)$  
   Uniform bubbles are formed

2. $2.4(N_c-1) < N_w < 16$  
   The bubble volume increases with the increase of $N_w$. 
3. \( N_w > 16 \)

The bubbles break down after detachment at the orifice.

where,

\[ N_w = \text{BoFr}^{0.5} \]

and Bo, Fr are the dimensionless Bond and Froude numbers respectively.

There are many theoretical and experimental studies concerning bubble formation at orifices (Kumar and Kuloor, 1970; Davidson and Schuler, 1960; Marmur and Rubin, 1973; Lanauze and Harris, 1973; Wraith, 1971; Marmur and Rubin, 1976; McCann and Prince, 1969). Davidson and Schuler (1960) have derived simple expressions for bubble volume under constant flow conditions in viscous and inviscid liquids. These equations are functions of gas rate and viscosity of continuous phase. The contradiction in the existing data, regarding the role of viscosity in bubble formation has been discussed by Kumar and Kuloor (1970). Generally the effect of viscosity is large at high gas flow rates (Kumar and Kuloor, 1970; Ruff, 1972; Wraith, 1971; Ramakrishnan, 1969) and bubble volume increases with viscosity of continuous phase. The same trend has been found for the case of constant pressure conditions (Kumar and Kuloor, 1970; Davidson and Schuler, 1960; Satyanarayan et al., 1969). Gas flow rate controls the mechanism of bubble formation. At low gas flow rates, separate bubbles are formed (Schugerl et al., 1977). At intermediate gas flow rates, bubble size is controlled by the instability of the jet interface (Meister and Scheele, 1967). At high gas
flow rates bubble size is controlled by interfacial forces and dynamic pressure forces of the local turbulence (Hinze, 1955).

Interfacial tension affects bubble volume at low flow rates and has a negligible effect at high flow rates (Kumar and Kuloor, 1970). However, a more detailed analysis takes into consideration the effect of continuous phase viscosity and interfacial tension (Satyanarayan, 1969). Another significant factor influencing bubble volume is the orifice diameter. At low gas rates, the effect of orifice size is such that the volume of the bubble increases with orifice size. At higher flow rates bubble volumes for various orifice diameters tend to join in a single curve (Kumar and Kuloor, 1970; Ramakrishnan, 1969).

Other parameters affecting bubble generation include: orifice characteristics, chamber volume, submergence, liquid density, gas physical properties, gas flow rate and continuous phase viscosity. Reported data are inconsistent and reflect a lack of appreciation by various investigators for the effects of chamber volume, the types of flows and the interactions of several variables considered (Clift et al., 1978). Because of these contradictions in the reported data bubble sizes are measured directly in the present work using a photographic technique.

2.2. Terminal Velocities of Bubbles

Bubbles/drops rise or fall freely in infinite media under the influence of
various forces e.g. gravitational, inertial, buoyancy and drag forces (Clift et al., 1978). Shape is controlled by the drag force which in turn determines the terminal velocity. For Reynolds numbers less than unity the inertia forces are negligible in comparison with viscous and interfacial tension forces. In this case, bubbles or drops are closely approximated by spheres. The terminal velocity and drag coefficient have been found analytically by Hadamard (1911) and Rybczynski (1911) who independently solved the Navier-Stokes equations. At high Reynolds numbers, spherical bubbles and drops are deformed to an ellipsoidal shape. In this case, surface tension, contaminants and secondary motion imposed by the internal circulation of fluid particle play an important role in determining terminal velocities. Surface contaminants are inevitable and for many practical purposes one has to consider them. Grace et al. (1976) has correlated a substantial amount of data reported in the literature. Bubble/drop deformation is more pronounced as the Reynolds number is increased further. For Re>150, the rear is quite flat and the wake angle $\theta_w$ is given (Clift et al. 1978) by,

$$\theta_w = 50^\circ + 190 \exp(-0.62 Re^{0.4})$$

Davies and Taylor (1950) have derived a simple expression for the terminal velocity of spherical-cap fluid particles for relatively high Reynolds numbers. Their calculations were based on equating the pressure distribution around the particle (potential flow) and the pressure distribution on the surface of the particle (hydrostatic pressure). Their
correlation is as follows:

\[ U_T = \frac{2}{3} \left[ g \alpha (\rho_\text{l} - \rho_\text{g}) / \rho_\text{l} \right]^{1/2} \quad \text{for} \quad 40 < \text{Re} < 150 \]  \hspace{1cm} (2.1)

Where \( U_T \) is the terminal velocity of bubble and \( \alpha \) the equivalent radius of the bubble. For Reynolds numbers greater than 150 and spherical-cap fluid particles the terminal velocity can be calculated from the following correlation (Clift et al. 1978) which is similar to the Davies and Taylor equation.

\[ U_T = 0.711 \left[ g d_e (\rho_\text{l} - \rho_\text{g}) / \rho_\text{l} \right]^{1/2} \quad \text{for} \quad \text{Re} > 150 \]  \hspace{1cm} (2.2)

Where \( d_e \) is the equivalent bubble diameter. This correlation was derived from the geometrical similarity of a spherical-cap fluid particle with a wake angle of approximately 50\(^\circ\), once \( \text{Re} \) is greater than about 150.

All the above equations are valid for individual bubbles. In a laminar flow field, bubbles interact and the above equations cannot be applied to estimate bubble swarm velocities. However, Ueyama and Miyauchi (1979) calculated bubble swarm velocities in a turbulent flow field using the above expressions with the Sauter mean diameter \( d_{\text{vs}} \) instead of \( d_e \) and found good agreement with experimental data. Thus, in a turbulent flow field the bubbles have limited interactions. Each bubble rises independently.

Therefore, the slip velocity of a bubble swarm can be defined as:

\[ u_s = 0.711 \left[ g d_{\text{vs}} (\rho_\text{l} - \rho_\text{g}) / \rho_\text{l} \right]^{1/2} \quad \text{for} \quad \text{Re} > 1000 \]  \hspace{1cm} (2.3)

The bubble swarm velocity can easily be related to the superficial gas
velocity $u_g$ through a simple expression, which is discussed in the next section.

2.3. Gas-Liquid Systems

Gas-liquid bubble column reactors are used extensively in chemical industries because of their various advantages i.e., low investment cost, low energy requirements and effective mixing. Examples include slurry-phase hydrogenation, aerobic biochemical reaction and polymerization of olefins. In this section some of the hydrodynamic parameters associated with gas liquid systems are discussed. Gas holdup is one of the most important hydrodynamic parameters characterizing such systems. It has two applications. On the one hand, it gives the volume fraction of gas in multi-phase systems. On the other hand, the gas holdup in combination with the mean bubble diameter allows the estimation of interfacial area. The existing correlations for gas holdup in the literature are functions of superficial gas velocity, column diameter, orifice size and physical properties of continuous phase (Hughmark, 1967; Akita and Yoshida, 1973; Kumar et al., 1976; Hikita et al., 1980; Shah et al., 1982). Gas holdup can be calculated directly from bed expansion as,

$$
\varepsilon_g = \frac{(H - H_o)}{H_o}
$$

(2.4)

where $\varepsilon_g$ is the gas holdup, $H_o$ the initial height of liquid in the column in cm
and $H$ is the head pressure in cm of liquid in the column, or indirectly by the manometric method:

$$\varepsilon_g = \frac{dP}{dh}$$  \hspace{1cm} (2.5)

Where $dP$ is the differential pressure drop over a differential height $dh$ of the column. The reported data related to gas holdup promote a scatter due to the sensitivity of the holdup to the gas-liquid system and to impurities (Shah et al., 1982). Abou-El-Hassan (1979) correlated a substantial amount of data with an expression that includes the effect of apparent viscosity. The correlation may be applied to gas-liquid-solid and gas-liquid-liquid systems.

Bubble rise velocity and bubble size distribution have a direct impact on the performance of bubble columns. Photographic techniques have been used widely for the study of such systems because of their simplicity (Schugerl et al., 1977; Akita and Yoshida, 1974; Burchart and Deckwer, 1975). Different statistical distribution functions have been found for bubble size by various workers. Akita and Yoshida (1974) found a logarithmic normal probability function, Buchart and Deckert (1975) found two different bubble size distributions coexisting in a column and Todtenhaupt (1971) a normal probability function. Though the original bubble size distributions differ markedly, the volume-surface mean drop diameter or Sauter mean diameter $d_{vs}$ differ only slightly. The $d_{vs}$ has been defined by Mugele and Evans (1951) as:
\[ d_{vs} = \left( \sum n_i d_{pi}^3 / \sum n_i d_{pi}^2 \right) \]  \hspace{1cm} (2.6)

However, these mean sizes depend on coalescence and break-up of bubbles and obviously on the kind of flow in the column. The superficial gas velocity has been defined for the case of a bubble swarm by Sriram et al. (1977) as:

\[ u_g = \varepsilon_g \int_{d_b}^{\infty} f(d_b) u(d_b) dd_b \]  \hspace{1cm} (2.7)

By writing the above equation for the case of a discrete function, the following expression can be obtained.

\[ u_g = \varepsilon_g \frac{\sum n_i ((i-1)\Delta d_b < d_b < i\Delta d_b) u_i ((i-1)\Delta d_b < d_b < i\Delta d_b)}{\sum n_i ((i-1)\Delta d_b < d_b < i\Delta d_b)} \]  \hspace{1cm} (2.8)

At low superficial gas velocities the dispersed gas moves freely as discrete bubbles in the liquid. This regime is known as uniform bubbling regime and has been studied by Molerus and Kurtin (1985). As the superficial gas velocity increases, the bubbles become unstable and localized liquid circulation is imposed on the flow field. Coalescence and break-up of bubbles can take place and there seems to be no regularity. This is the transition flow regime from the bubbling regime to the turbulence...
regime and it has been explicitly treated by Maruyama et al. (1979). The turbulent flow regime which has been studied by Ueyama and Miyauchi (1979) is of practical importance. In this case, the circulation is clear in the column due to the existence of a phase rich in bubbles in the central plume of the column and a phase relatively lean in bubbles near the wall. Thus there exists a difference in buoyancy forces, which cause the circulation of the continuous phase. Provided that these conditions are met, isotropic turbulence theory can be applied once the length and velocity scales of the inertial subrange eddies are known.

2.4. Break-up and Coalescence of Drops in Turbulent Flow Fields-The Effect of Energy Dissipation

The intensity and structure of turbulence is of great importance, as it controls the fluid drop size and size distribution. A statistical description of the structure of turbulence originally proposed by Kolmogoroff (1941) provides a basis for describing fluid particle break-up and coalescence. Such phenomena play an important role in determining interfacial areas in hydrorefining processes (Shaw, 1985) and in other multiphase systems where the mass transfer rates must be optimized.

The simplest form of turbulence is that of isotropic turbulence and has been defined by the condition that all mean values of functions of the flow variables should be independent of rotation and reflection of the axes of
The turbulent eddies are divided into three categories. The energy-containing eddies which are of large wave numbers and depend on the characteristics of flow; The eddies of the inertial subrange which play the role of transferring the kinetic energy from the energy-dissipating eddies to the viscous eddies; and the energy-dissipating eddies, the smallest ones with respect to length scale, which are independent of the flow characteristics. The latter eddies depend on the energy lost from the large energy-containing eddies and once this energy rate is known their structure is determined. No matter how inhomogeneous the flow is, isotropic turbulent flow may be considered locally in the range of energy-dissipating eddies (Batchelor,1960). From dimensional reasoning, Kolmogoroff defined the length and velocity scales for the universal equilibrium range (inertial and viscous subranges) as:

\[ \eta = (\nu^3 / \varepsilon)^{1/4} \quad \text{and} \quad u = (\nu \varepsilon)^{1/4} \]

Where \( \eta \) is the length scale, \( u \) is the velocity scale, \( \nu \) is the kinematic viscosity of the continuous liquid phase and \( \varepsilon \) is the energy dissipation rate per unit mass of liquid. The length scale \( L \) of the energy-containing eddies is comparable to the physical dimensions of the system. If \( d_p \) is the diameter of a drop and

\[ \eta < d_p < L \]

then, the variation of dynamic pressure fluctuations in a distance equal to
\( d_\text{p} \) is strong enough to cause drop break-up. The maximum diameter of a drop, stable against break-up in turbulent flow fields is,

\[
d_{\text{max}} \sim (\sigma/\rho)^{3/5} \epsilon^{-2/5}
\]

(2.9)

and has been defined by Hinze (1955). However, there also exists a minimum drop size, stable against coalescence in turbulent flow fields which has been derived by Thomas (1981) as:

\[
d_{\text{min}} \sim 2.4 \left( \frac{\sigma^2 h^2}{\mu \rho \epsilon} \right)^{1/4}
\]

(2.10)

As \( d_{\text{max}} \) is over an order of magnitude greater than \( d_{\text{min}} \), a broad range of drop sizes is stable in a turbulent field. This broad range narrows as the energy dissipation rate is increased. There exists a critical value of the energy dissipation \( \epsilon_o \) where the \( d_{\text{min}} \) and \( d_{\text{max}} \) are theoretically the same (Thomas, 1981). Therefore, if \( \epsilon < \epsilon_o \) drops with diameters lying between \( d_{\text{max}} \) and \( d_{\text{min}} \) are too large to coalesce and also too small to be broken up. So, the final size distribution is controlled by the magnitude of the energy dissipation \( \epsilon \).

In agitated vessels, the energy dissipation rate, \( \epsilon \), is related to the impeller speed and vessel geometry:

\[
\epsilon = C D^2 N^3
\]

(2.11)
Where $D$ is the diameter of the impeller, $N$ is the number of revolutions of the impeller per unit time and $C$ is a proportionality constant strongly depended on impeller type and size.

In tubular reactors for gas-liquid and gas-liquid-liquid systems the energy dissipation rate can be found approximately by ignoring the gas holdup and density of the gas

$$\varepsilon = u_g g$$  \hspace{1cm} (2.12)

A more accurate expression for calculating the energy dissipation rate in tubular reactors is derived in APPENDIX A.

### 2.5. Dispersion Phenomena in Agitated Vessels

Many operations in chemical engineering require the contact of two liquid phases. One of the most common methods of bringing about the contact of two liquid phases is to disperse drops of one within the other by mechanical agitation. The interfacial area is affected by the physical and chemical characteristics of the system and mechanical features of the equipment. These characteristics as well as their impact on the Sauter mean drop diameter in agitated vessels are presented in this section.

A number of investigators have presented correlations that relate drop sizes in agitated vessels to mixing parameters and physical properties of the system (Weinstein and Treybal, 1973; Coulaloglou and Tavlarides, 1976; Nishikawa et al., 1987; Stamatoudis and Tavlarides, 1985). Drop break-up
has been attributed to turbulent pressure fluctuations as discussed above. In agitated vessels, two distinct operating regimes have been observed. The Sauter mean drop diameter $d_{vs}$ is controlled by different mechanisms in each of these regimes. At low impeller speeds drop break-up controls the Sauter mean diameter $d_{vs}$ which is correlated as a function of energy dissipation, $\varepsilon$, to the power $-0.4$ (Weinstein and Treybal, 1973; Nishikawa, 1987). At high impeller speeds the dominant factor determining drop size is not break-up but the prevention of coalescence by the action of turbulence as discussed by Shinnar (1957, 1961) and Shinnar and Church (1960). This regime is known as coalescence prevention regime and the Sauter mean drop diameter was found to be proportional to the energy dissipation to the power $-0.25$ (Nishikawa, 1987).

In agitated vessels, local mean drop diameter measurements have indicated that there exists a non-uniform drop size distribution. Smaller drops have been found to exist near the impeller (Weinstein and Treybal, 1973; Coulaloglou and Tavlarides, 1976). This is due to the fact that the energy is not dissipated uniformly over the volume of the vessel.

The effect of dispersed liquid volume fraction, $\phi$, depends on the operating regime and its role becomes more significant in the coalescence-prevention regime (Nishikawa, 1987). Generally, an increase of $\phi$ gives a larger Sauter mean drop diameter. However, it has to be pointed
out that the volume fraction, $\phi$, has no effect on the Sauter mean diameter for lean dispersions less than 0.5% (Narshimhan, 1980). The effect of relative viscosity, $\mu_d/\mu_c$, between dispersed liquid phase and continuous liquid phase has been found to have a considerable impact on the Sauter mean drop diameter. Nishikawa (1987) reported a power dependence of $d_{vs}$ on $\mu_d/\mu_c$ ranging from 0.125 to 0.25. Calderbank (1967) proposed a number of equations for $d_{vs}$ including the term $(\mu_d/\mu_c)^{0.25}$. Hinze (1955) in his classical paper points out that the greater the deviation of relative viscosity is from unity the greater the resistance to drop break-up.

Unsteady-state drop size distributions in agitated vessels have been studied by Narsimhan et al. (1979, 1980) and Ramkrishna (1974). The Sauter mean diameter changes relatively faster during the early stages of the operation and the rate of change may be correlated as an exponential function. Moreover, a change in the impeller speed also promotes a change of Sauter mean diameter in a comparatively small amount of time.

Nishikawa (1987) has summarized in the following equations the impact of physical properties of liquid phases and operating conditions on Sauter mean diameter:

$$d_{vs} = 0.0371 \, \varepsilon^{-1/4} \, (d/D)^{3/4} \, (1+3.5 \, \phi^{3/4})(\mu_d/\mu_c)^{13/40} \, (\sigma/\rho)^{3/8} \quad (2.13)$$

(coalescence-prevention regime)
\[ d_{vs} = 0.1050 \varepsilon^{2/5} (d/D)^{6/5} (1+2.5 \varphi^{2/3})(\mu_d/\mu_c)^{13/40} (\sigma/\rho)^{3/8} \tag{2.14} \]

(break-up regime)

Where \( d \) is the diameter of the impeller and \( D \) is the diameter of the vessel.

A detailed analysis reveals that the average power input per unit mass of liquid is not a satisfactory method of stirred vessel scale-up. Power is employed more advantageously by using a large impeller rotating at low speed rather than a short impeller at high speed. As it is shown from the above correlation there is a strong dependence of Sauter mean diameter on the geometry of the vessel. Therefore, it is an unreliable method to use Equations 2.13 and 2.14 for other geometrical arrangements.

2.6. Objectives

As it is shown in the literature review there is little information concerning gas-liquid-liquid systems. Therefore, there is a need for an extensive and detailed study of such systems. A number of different characteristics associated with phase interactions and hydrodynamics of gas-liquid-liquid systems will be addressed. The present investigation focusses on:

1. **Gas Flux and Physical Properties of Dispersed Phase Liquid**: since gas flux is directly related to energy dissipation rate, it is expected to play an important role in determining the mean drop size. Physical properties of dispersed phase liquid, including interfacial tension, viscosity and density,
are also expected to have a significant impact on the steady state mean drop size and drop size distribution.

2. **Initial Drop and Bubble Size:** different bubble sizes impose different shear fields which may result in different interfacial areas as is discussed for the case of agitated vessels (Section 2.5).

3. **Unsteady State Drop Size Distribution:** the transient drop size distribution is studied so that interfacial area can be modelled for reactions where steady state drop size distributions are not obtained.

4. **Maximum and Minimum Stable Drop Sizes:** drops stable against break-up and coalescence set bounds on the size spectrum of drops in a reactor. Formulae applicable to tubular reactors are derived. Such formulae are very useful for correlating the Sauter mean diameter.

5. **Comparison between Tubular Reactors and Agitated Vessels:** agitated vessels and tubular reactors are common phase-contacting devices. A comparison between them based on the rate of energy dissipation reveals their relative performance.
PART II

EXPERIMENTAL RESULTS
3.0. EXPERIMENTAL

3.1. Description of the Experimental Apparatus

All the experiments were carried out in a column with a rectangular cross-section of 15 X 30 cm in order to simulate an axial slice of a tubular reactor. A general schematic of the apparatus is shown in Figure 1. A detailed sketch of the column and gas chamber is shown in Figure 2. The rectangular cross-section was chosen to facilitate the photographic technique used in the present work. The thickness of the column is relatively large for such simulation. However, this was chosen to avoid wall effects. The column, 100 cm in height, was constructed of 0.635 cm thick sheets of perspex. Five taps of 1.27 cm diameter were drilled along the height of the column at 20 cm intervals, the lowest being 5 cm above the bottom plate so that the pressure drop could be measured. Compression fittings were used for the connections in order to avoid leaks. Five holes were drilled in the bottom plate of the column and fitted with interchangeable nozzles (0.15, 0.25, 0.35 and 0.5 cm diameter). The distance between the nozzles was 5 cm in order to avoid interactions between neighbouring bubbles, as they are generated.

The holes in the bottom plate linked the reactor to a gas chamber with a rectangular cross-section of 30 X 40 cm and height of 40 cm. This tank was made up of 0.635 cm thick sheets of perspex and had a top plate constructed of 1.27 cm thick sheet of perspex. A 25 X 2 cm slot and four capillaries
were drilled into the top plate to promote uniform gas distribution. The volume of the chamber was varied by filling it with water. This feature in conjunction with the various size nozzles permitted bubble generation at constant pressure, intermediate and constant flow conditions as discussed in section 2.2. A pressure relief valve set at 1.3 atm was used to prevent overpressuring the apparatus.

Two different rotameters were used to control the gas flow at low and high gas rates. At low gas rates, a 600 series rotameter of Matheson with a 15 cm tube (7632T) was used, equipped with a glass float. The typical flow range was from 6 cm$^3$/sec up to 150 cm$^3$/sec. A high accuracy needle valve was placed at the inlet of the rotameter to get the downstream pressure close to atmospheric. The upstream pressure was less than 138 kPa, as specified by the rotameter manufacturer. At high gas rates a Fischer & Porter rotameter was used with a typical flow range from 100 cm$^3$/sec up to 1750 cm$^3$/sec. Calibration curves for both rotameters are shown in Figure 3.

Cylinder air was used to generate bubbles and a two-stage brass regulator Matheson (Model 8) was used to control the pressure. The delivery pressure gauge could be varied in the range of 0-690 kPa. A copper-pipe 6 cm I.D. and 60 cm long was used to saturate the air with water in order to avoid mass transfer in the column. The saturator was filled with the same liquid as the continuous liquid phase.
U-glass-tube manometers 0.5 cm I.D. were used for the following purposes:
- To measure the pressure fluctuations under the gas-distributor plate because of the bubble generation.
- To measure the pressure in the gas chamber.
- To measure the pressure at the outlet rotameter.
- To measure the pressure along the column.

A single experimental setup was devised to generate various drop sizes. The device comprised a 2 lit volume flask as a tank for the dispersed phase liquid, three liquid distributors with orifice sizes (0.32, 0.16, 0.8 cm) and a valve to control the liquid flow rate.

### 3.2. Experimental procedure

The reactor was operated in batch mode with respect to both continuous and dispersed phase liquids and in continuous mode with respect to gas phase. The physical properties of various G-L-L systems used for this investigation are listed in Table 1. Sources and purities of these chemicals are tabulated in Table 2. Viscosity was measured using an Ostwald viscometer and interfacial tension was measured using a ring intertensiometer. During a typical experiment, the column was filled with distilled water to a depth of 85 cm and air was passed through the nozzles at a flow rate ranging from 45 to 1750 cm³/sec. A perforated plate was placed at the top of the reactor and about 5 cm below the water level in
order to minimize free surface effects. The dispersed phase liquid was introduced from the top of the reactor through various size nozzle distributors (0.32, 0.16 or 0.1 cm). The initial drop size was greater than 2000 µm. Drop and bubble diameters were determined photographically using a Nikon camera (Shutter speed 1/1000th sec) in conjunction with an image analyzer (Bausch and Lomb, Omnicon-3000 with NOVA-4 data general). At low gas fluxes, photographs were taken while the column was operating. However, at higher gas fluxes photographs were taken only at the end of an experiment. To study the dependence of drop size on time at high gas fluxes the following procedure was used. After stopping the air supply at fixed time intervals, photographs were taken immediately and the experiment was repeated all over again in order for photographs to be taken at the next fixed time interval. Preliminary experiments showed that the mean drop diameter remained virtually constant after 15 minutes of operation. A typical image analysis was based on the size analysis of 1000 plus drops. Size analysis of approximately 500 drops gave similar results. Correction factors had to be used due to geometrical magnifications involved in photographic techniques. The repeatability of the above described experimental procedure is within 5% as shown in Table 3.
Figure 1: Schematic diagram of experimental apparatus
Figure 2: Detailed sketch of bubble column and gas chamber
Figure 3: Calibration curves for the rotameters: a) Matheson (7632T) 
b) Fischer & Porter
### TABLE 1. Physical properties of Gas-liquid-liquid systems.

<table>
<thead>
<tr>
<th>Systems</th>
<th>$\mu_d$, poise</th>
<th>$\sigma$, dyne/cm</th>
<th>$\rho_d$, g/cm$^3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. Linseed oil (96%/v)+TCE (4%/v)</td>
<td>0.25</td>
<td>6.1</td>
<td>0.95</td>
</tr>
<tr>
<td>Water-Air</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>B. Linseed oil (85%/v)+TCE (15%/v)</td>
<td>0.17</td>
<td>6.2</td>
<td>1.00</td>
</tr>
<tr>
<td>Water-Air</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C. Linseed oil (82%/v)+TCE (18%/v)</td>
<td>0.15</td>
<td>6.4</td>
<td>1.02</td>
</tr>
<tr>
<td>Water-Air</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>D. Linseed oil (78%/v)+TCE (22%/v)</td>
<td>0.13</td>
<td>6.8</td>
<td>1.04</td>
</tr>
<tr>
<td>Water-Air</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E. Linseed oil (61%/v)+TCE (39%/v)</td>
<td>0.03</td>
<td>10.2</td>
<td>1.13</td>
</tr>
<tr>
<td>Water-Air</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F. Cottonseed oil (83%/v)+TCE (17%/v)</td>
<td>0.19</td>
<td>23.6</td>
<td>1.00</td>
</tr>
<tr>
<td>Water-Air</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>G. Dibenzyl Ether-Water-Air</td>
<td>0.05</td>
<td>10.2</td>
<td>1.04</td>
</tr>
</tbody>
</table>

Systems B, F and G are referred to as the linseed oil, cottonseed oil and dibenzyl ether systems respectively in the text. Systems A, C, D and E have only been used to quantify the impact of relative density and viscosity.
TABLE 2. Sources and Purities of Chemicals

<table>
<thead>
<tr>
<th>Chemicals</th>
<th>Purity (%)</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air</td>
<td>99</td>
<td>CANOX</td>
</tr>
<tr>
<td>Water</td>
<td>High purity</td>
<td>U. of TORONTO</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Dept. of Chem. Eng.</td>
</tr>
<tr>
<td>Linseed oil</td>
<td>-</td>
<td>RECOCHEM</td>
</tr>
<tr>
<td>Cottonseed oil</td>
<td>For laboratory use</td>
<td>FISCHER Scientific Company</td>
</tr>
<tr>
<td>Dibenzyl ether</td>
<td>98</td>
<td>FLUKA AG (Switzerland)</td>
</tr>
<tr>
<td>Trichloroethylene</td>
<td>99</td>
<td>BDH Chemicals</td>
</tr>
</tbody>
</table>
### TABLE 3. Repeatability of the Procedure.

<table>
<thead>
<tr>
<th>System</th>
<th>$D_{\text{max}}$, $\mu$m</th>
<th>$D_{\text{vs}}$, $\mu$m</th>
<th>$D_{\text{min}}$, $\mu$m</th>
<th>$u_g$, cm sec$^{-1}$</th>
<th>$\phi(%)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>B</td>
<td>926</td>
<td>465</td>
<td>76</td>
<td>0.1</td>
<td>0.45</td>
</tr>
<tr>
<td></td>
<td>945</td>
<td>472</td>
<td>78</td>
<td>0.1</td>
<td>0.45</td>
</tr>
<tr>
<td>A</td>
<td>1051</td>
<td>520</td>
<td>88</td>
<td>0.1</td>
<td>0.45</td>
</tr>
<tr>
<td></td>
<td>1055</td>
<td>505</td>
<td>83</td>
<td>0.1</td>
<td>0.45</td>
</tr>
<tr>
<td>F</td>
<td>521</td>
<td>262</td>
<td>43</td>
<td>0.4</td>
<td>0.45</td>
</tr>
<tr>
<td></td>
<td>532</td>
<td>269</td>
<td>49</td>
<td>0.4</td>
<td>0.45</td>
</tr>
<tr>
<td>G</td>
<td>545</td>
<td>274</td>
<td>45</td>
<td>0.2</td>
<td>0.45</td>
</tr>
<tr>
<td></td>
<td>529</td>
<td>279</td>
<td>49</td>
<td>0.2</td>
<td>0.45</td>
</tr>
</tbody>
</table>

The maximum deviations which can be observed from the above listed results vary from 2% to 5%.
PART III

RESULTS AND DISCUSSION
4.0. RESULTS

In this section the results obtained in the present investigation are presented in the form of Tables and Figures. These results are discussed in chapter 5.

Table 4 shows the maximum, minimum and Sauter mean diameters of dispersed phase drops for the bulk of experiments carried out in the present investigation. The operating conditions and systems used are also shown.

Figure 4 shows the gas holdup in gas-liquid-liquid tubular reactors. On the same Figure the generalized correlation of Abou-El-Hassan (1979) is shown which is valid for two and three phase systems. Kato's correlation (1984) is also sketched on the same Figure for gas-liquid-liquid tubular reactors operating in a continuous fashion with respect to all phases. Figures 5 and 6 are typical examples of size distributions and cumulative drop size distributions of dispersed phase droplets respectively for a variety of operating conditions and systems.

4.1. Unsteady State Drop Size Distributions

Figures 7, 8 and 9 illustrate unsteady state cumulative volume distributions for various systems and operating conditions used in this investigation. Figures 10, 11 and 12 show the effect of superficial gas velocity, interfacial tension and viscosity on the rate at which steady state
is reached in three phase tubular reactors. Figures 13 and 14 are plots of drop diameter against time for various cumulative volume values. These Figures are based on Figures 7, 8 and 9 and are very helpful in identifying the similarity variable (Section 5.3.2). Figures 15 and 16 show the significance of the similarity variable. Finally Figure 17 is a comparison between the rate at which steady state drop size distributions are approached in tubular reactors and agitated vessels.

4.2. Steady State Drop Size Distributions

Figure 18 shows the effect of superficial gas velocity on the Sauter mean diameter of dispersed liquid drops for three different systems. The results are correlated (Equation 5.16) and a parity plot for this correlation is shown in Figure 19. The functional dependence of maximum and minimum stable drop sizes against break-up and coalescence respectively are shown on Figures 20 and 21 for two different systems. The various slopes appearing on Figures 20 and 21, calculated by using least square method, are tabulated in Table 5. Figures 22, 23, 24 and 25 show the effect of energy dissipation rate on the breadth of the drop size distribution for various systems.

Figure 26 shows the effect of relative viscosity on the steady state drop size. Figures 27 and 28 show the effect of dispersed phase volume fraction on the steady state Sauter mean diameter for two different
systems. In Table 6, the effect of density difference between the two liquid phases is shown.

Finally, Figure 29 shows a comparison of the relative performance of tubular reactors (experimental data) and agitated vessels (Equations 2.13 and 2.14) for three different systems:

- a system with high interfacial tension and high relative viscosity
- a system with low interfacial tension and high relative viscosity
- a system with low interfacial tension and low relative viscosity.
### TABLE 4. Steady state Max, Min and Sauter Mean Diameters.

<table>
<thead>
<tr>
<th>System</th>
<th>(D_{\text{max}}, \mu\text{m})</th>
<th>(D_{\text{vs}}, \mu\text{m})</th>
<th>(D_{\text{min}}, \mu\text{m})</th>
<th>(u_g, \text{ cm sec}^{-1})</th>
<th>(\phi(%))</th>
</tr>
</thead>
<tbody>
<tr>
<td>B</td>
<td>926</td>
<td>465</td>
<td>76</td>
<td>0.1</td>
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<tr>
<td></td>
<td>490</td>
<td>246</td>
<td>40</td>
<td>0.4</td>
<td>1.00</td>
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TABLE 5. Slopes of Various Systems in Both Flow Regimes

<table>
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<tr>
<th>System</th>
<th>Slopes in Low Gas Flux Flow Regime</th>
<th>Slopes in High Gas Flux Flow Regime</th>
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<tr>
<td></td>
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</tr>
<tr>
<td>F</td>
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<td>0.55</td>
</tr>
<tr>
<td>G</td>
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</table>
TABLE 6. Effect of relative density $\rho_d/\rho_c$

<table>
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<tr>
<th>System</th>
<th>$\rho_d/\rho_c$</th>
<th>$\mu_d/\mu_c$</th>
<th>$\sigma$, dyne/cm</th>
<th>$u_g$, cm/sec</th>
<th>$d_{vs}$, $\mu$m</th>
<th>$d_{vs, cor.}$, $\mu$m</th>
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<td>520</td>
<td>441</td>
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<td>1.00</td>
<td>17</td>
<td>6.2</td>
<td>0.1</td>
<td>465</td>
<td>434</td>
</tr>
<tr>
<td>C</td>
<td>1.02</td>
<td>15</td>
<td>6.4</td>
<td>0.1</td>
<td>459</td>
<td>442</td>
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<td>D</td>
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<td>13</td>
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<td>6.2</td>
<td>0.2</td>
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<tr>
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<td>10.2</td>
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<tr>
<td>B</td>
<td>1.00</td>
<td>17</td>
<td>6.2</td>
<td>0.4</td>
<td>264</td>
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<tr>
<td>D</td>
<td>1.04</td>
<td>13</td>
<td>6.8</td>
<td>0.4</td>
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<tr>
<td>E</td>
<td>1.13</td>
<td>3</td>
<td>10.2</td>
<td>0.4</td>
<td>202</td>
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</table>

$d_{vs, cor.}$ is the corrected value of Sauter mean diameter accounting for the different relative viscosity $\mu_d/\mu_c$ of various systems. Basis is taken $\mu_d/\mu_c=13$. 
Figure 4: Gas holdup in three phase tubular reactors
Figure 5: Sample size Distributions for dispersed phase drops
Figure 6: Sample cumulative drop size distributions
Figure 7: Unsteady state cumulative volume distributions for the dibenzyl ether system ($u_g = 0.1 \text{ cm/sec}$)
Figure 8: Unsteady state cumulative volume distributions for the cottonseed oil system ($u_g=0.4$ cm/sec)
Figure 9: Unsteady state cumulative volume distribution for the linseed oil system ($u_g = 2$ cm/sec)
Figure 10: The effect of superficial gas velocity, $u_g$, on the unsteady state mean drop size (linseed oil system)
Figure 11: The effect of interfacial tension, \( \sigma \), on the unsteady state mean drop size \( (u_g=0.1 \text{ cm/sec}) \)
Figure 12: The effect of relative viscosity, $\mu_o/\mu_c$, on the unsteady state mean drop size ($u_g = 0.1 \text{ cm/sec}$)
Figure 13: The functional dependence of the transition probability function $\Gamma(v)$ on drop size (dibenzyl ether system, $u_g=0.1$ cm/sec)
Figure 14: The functional dependence of the transition probability function $I(v)$ on drop size (linseed oil system, $u_g=2$ cm/sec)
Figure 15: The similarity of the unsteady state cumulative volume size distributions (dibenzyl ether system, $u_g=0.1$ cm/sec)
Figure 15: The similarity of the unsteady state cumulative volume size distributions (linseed oil system, $u_g=2$ cm/sec)

Figure 16: The similarity of the unsteady state cumulative volume size distributions (linseed oil system, $u_g=2$ cm/sec)
Figure 17: A comparison between the rate at which steady state drop size distributions are approached in tubular reactors and agitated vessels.
Figure 18: The effect of superficial gas velocity on the steady state Sauter mean diameter of dispersed liquid drops
Figure 19: Parity plot for Equation 5.16
Figure 20: The dependence of maximum and minimum drop sizes on the energy dissipation rate (cottonseed oil system)
Figure 21: The dependence of maximum and minimum drop sizes on the energy dissipation rate (linseed oil system)
Figure 22: The effect of energy dissipation rate on the breadth of drop size distributions for the linseed oil system.
Figure 23: The effect of energy dissipation rate on the breadth of drop size distributions for the cottonseed oil system
Figure 24: The effect of energy dissipation rate on the size spectrum of dispersed phase drops (cottonseed oil system)
Figure 25: The effect of energy dissipation rate on the cumulative size spectrum of dispersed phase drops (linseed oil system)
Figure 26: The effect of relative viscosity, $\mu_d/\mu_c$, on the steady state Sauter mean drop size
Figure 27: The effect of volume fraction, \( \phi \), of the dispersed phase on the steady state Sauter mean diameter (linseed oil system)
Figure 28: The effect of volume fraction, $\phi$, of dispersed phase on the steady state Sauter mean diameter (cottonseed oil system)
Figure 29: A comparison of the performance of tubular reactors (experimental data) and agitated vessels (Equations 2.13 and 2.14 with d/D=0.5) at: a) high relative viscosity and low interfacial tension, b) high relative viscosity and high interfacial tension, c) low relative viscosity and low interfacial tension.
5.0. DISCUSSION

The results obtained in this investigation are presented in chapter 4 in the form of Tables and Figures. In this chapter these results are discussed and some characteristics not obviously extracted from the Figures are presented and explained. Gas holdup as a hydrodynamic parameter characterizing the operation of gas-liquid-liquid tubular reactors is discussed first. Then, unsteady state drop size distributions and unsteady state mean drop size are presented as fundamental variables in characterizing the time dependence and transition of drop breakage in these processes. Subsequently, the importance of Sauter mean drop diameter in estimating the interfacial area as well as the size spectrum of stable drops, are discussed and correlated. The impact of physical properties of the dispersed liquid phase on Sauter mean diameter is assessed and quantified in conjunction with the impact of superficial gas velocity. Finally, the results of this investigation are compared with stirred tank analogues.

5.1. Gas Holdup

Figure 4 shows the relationship between gas holdup and superficial gas velocity in the reactor. The mean gas holdup $\varepsilon_g$ is correlated as:

$$\varepsilon_g = 0.375 \, u_g^{0.575}$$  \hspace{1cm} (5.1)

where $u_g$ is expressed in the units of cm/s. Shah et al. (1982) in a review
paper pointed out that the gas holdup depends mainly on superficial gas velocity and is often very sensitive to the kind of liquid media for gas-liquid systems. The number of correlations proposed indicates that no single unified equation is available. However, the exponents of $u_g$ in most of the proposed correlations particularly for churn turbulent flow are very close to 0.5 (Ueyama and Miyauchi, 1979).

As seen in Figure 4, the gas holdup in the range of 0.1-4 cm/s for linseed oil and cottonseed oil systems are similar indicating that the physical properties of the dispersed liquid phase do not have a significant effect. These values are comparable to values given by Abou-El-Hassan (1979), who devised a generalized correlation for gas holdup in gas-liquid and gas-liquid-solid bubble columns. The presence of a dispersed phase leads to a reduction in gas holdup vis-a-vis corresponding gas-liquid system. Kato et al. (1972) and Nakamura (1978) observed a similar reduction in gas holdup in bubble columns containing suspended solid particles. Ying et al. (1980) and Vasalos et al. (1980) investigated the effect of solids concentration and concluded that an increase in solid concentration generally decreases the gas holdup. This reduction in gas holdup has been attributed to an increase in the apparent viscosity of the bulk liquid resulting from the presence of a dispersed phase (Abou-El-Hassan, 1979) and would appear to be independent of the physical properties of the dispersed phase. Results obtained as part of this study are consistent with this finding as indicated in Figure 4. Kato et al. (1984)
found that gas holdup in gas-liquid-liquid systems tended to be 10-15% lower than values encountered in gas-liquid systems. Their correlation (dashed line in Figure 4) severely underestimates gas holdup in gas-liquid-liquid columns at low gas fluxes and overestimates it at high gas fluxes. However, the operation of their column was continuous with respect to all three phases. In batchwise operation the liquid in the wake of bubbles moves faster than the continuous liquid phase. As a result, the average velocity of the bulk liquid phase decreases. Since gas holdup is increased with the bulk liquid velocity, a slightly higher holdup is observed in cocurrent continuous operation.

5.2. Drop Size Distribution-Steady state

Typical steady state drop size distributions as well as steady state cumulative drop size distributions are illustrated in Figures 5 and 6 respectively. These distributions based on more than 1000 drops per distribution are approximately Gaussian as shown in Figure 5. However, the steady state size distribution is slightly skewed towards the smaller drop sizes. This is understandable as only a very small number of drops of size greater than $d_{\text{max}}$ can exist at equilibrium. Also due to uniform breakage of droplets over the volume of the reactor, one may expect a broad spectrum of drops less than $d_{\text{max}}$ and a comparatively small number of drops of size less than $d_{\text{min}}$. Similar distributions have been reported for kerosene-water
dispersions (Stamatoudis and Tavlarides, 1985) and for a variety of other liquid-liquid systems (Narsimhan et al., 1979) in agitated vessels.

The transient drop size distribution is modelled in section 5.3. It can also be used for determining the steady state drop size distribution, once the time to reach steady state has been identified (Hatzikiriakos et al., 1988c).

5.3. Drop Size Distribution-Unsteady state

The design of gas-liquid-liquid tubular reactors requires not only a knowledge of steady state drop size distribution, but also the dynamic characteristics of drop break-up and coalescence. The dynamic character of drop break-up and coalescence has been recognized by various workers (Coulaloglou and Tavlarides, 1977; Shah and Ramkrishna, 1973 and Narsimhan et al., 1980). These workers have approached the problem of prediction of drop size distributions by using population balances. However, such approaches cannot predict breakage and coalescence rates and the dynamic interrelationship between them. In this section an attempt is made to characterize the dynamic features of drop breakage in gas-liquid-liquid tubular reactors for lean dispersions. Following Ramkrishna (1973), the probability of drop break-up is quantified and suggestions for evaluating the cumulative drop size distribution function are given. Finally, a comparison between the rate at which steady state drop size distributions are approached in tubular reactors and agitated vessels is performed.
5.3.1. Dynamics of Drop Breakage in Tubular Reactors

The effect of time on drop break-up was studied for lean dispersions (volume fraction=0.0045). Therefore, the effect of drop coalescence may be considered negligible. Three different gas fluxes ($u_g=0.1, 0.4 \text{ and } 2 \text{ cm/sec}$) and three different systems (system B, F and G. Table 1) were chosen. During a typical experiment drop size distributions were obtained after 3, 6,10,15, 30 and 45 minutes. Initially, it was identified that at low gas fluxes steady state was reached in less than 30 minutes. Thus, subsequent drop size distributions were only obtained after 3, 6,10,15 and 30 minutes.

In Figures 7, 8 and 9 the cumulative volume distributions of dibenzyl ether, cottonseed oil and linseed oil systems at low, immediate and high gas fluxes respectively are plotted. Time is a parameter. It can be seen that large drops present initially are rapidly broken up and a steady state size distribution profile is obtained in less than 30 minutes. From the same Figures, it may also be observed that as time increases the breadth of drop size distribution narrows and the probability density steepens. Figure 10 shows the impact of superficial gas velocity on the rate at which steady state is reached. For relatively large times ($t>3 \text{ min}$) the rates at different gas fluxes are parallel, indicating that no effect exists. Similar observations may be inferred from Figures 11 and 12 where the effects of interfacial tension and relative viscosity are shown. The rates are parallel for times greater than 3 minutes. Therefore, the physical properties of dispersed phase and superficial gas velocities do not have significant effect
on the rate at which steady state is reached. The reduction of mean drop diameter with respect to time for the three systems under investigation is correlated as:

\[ d_{vs,t} = d_{vs} (1 + 1.6 \exp(-t/5)) \]  

(5.2)

where \( d_{vs,t} \) is the mean drop size at time \( t \), \( d_{vs} \) is the Sauter mean diameter at steady state and \( t \) is time in minutes. Equation 5.2 is represented by the solid line on Figure 17. As it may be seen, it fits the experimental data well. For \( t=0 \), the above Equation becomes \( d_{vs,t} = 2.6d_{vs} \). This suggests that at times less than 3 minutes the rate of break-up may depend on the initial drop size, physical properties of the system and superficial gas velocity. However, for \( t>3 \) min a similarity in the rate of drop breakage exists for the systems under investigation.

5.3.2. Similarity and Probability of Drop Breakage

In section 5.3.1 it was concluded that dispersed phase properties and superficial gas velocity do not have a significant effect on the rate at which steady state is reached. This suggests that a similarity in drop breakage should exist with respect to time. In the present section this similarity is identified and subsequently quantified. It is also demonstrated that measurements of cumulative volume distributions of drop sizes at various instants provide all the required information for prediction of unsteady state drop size distributions. The knowledge of drop size distribution at
every instant is of great importance because of the simultaneous nature of drop breakage and transport processes.

If the volume fraction of drops with a volume less than $v$ at time $t$ is denoted by $F(v,t)$ which is a cumulative distribution function, then this function satisfies the following conditions.

\[ F(v,t) = 0 \text{ when } v \leq 0 ; \quad F(v,t) = 1 \text{ when } v \to \infty \]

If $\Gamma(v)$ is the transition probability function then $\Gamma(v) \, dt$ represents the probability that a drop of volume $v$ breaks in the time interval $(t,t+dt)$. The volume fraction of daughter drops with volume less than $v$ formed from breakage of a drop of volume $v'$ represented by the function $G(v,v')$, has the following properties.

\[ G(v,v') = 0 \text{ when } v \leq 0 ; \quad G(v,v') = 0 \text{ when } v \geq v' \]

Ramkrishna (1973) showed that for a batch operation with lean dispersions, Equation 5.3 can be written,

\[
\frac{\partial F}{\partial t} = \int \frac{\Gamma(v') \, G(v,v') \, dF(v',t)}{F(v,t)}
\]

which can be rewritten as

\[ \frac{\partial F}{\partial t} = \int \frac{\Gamma(v') \, G(v,v') \, dF(v',t)}{F(v,t)} \]

(5.3)
\[
\frac{\partial F}{\partial t} = \int_{v}^{\infty} \Gamma(v') G(v,v') \frac{\partial F}{\partial v'} (v',t) \, dv'
\]  
(5.4)

where \( \frac{\partial F}{\partial v} \) is the density function. At this point \( \Gamma(v') \) and \( G(v,v') \) must be known functions in order to quantify the cumulative distribution function \( F(v,t) \). Ramkrishna (1973) made two assumptions for drop breakage of lean dispersions in agitated vessels. The breakage probability function was assumed to be of the form

\[
\Gamma(v) = K v^n
\]  
(5.5)

where \( K \) and \( n \) are constants to be identified from experimental measurements. The function \( G(v,v') \) was also assumed to be of the form

\[
G(v,v') = g(v/v') \quad 0 < v/v' < 1
\]  
(5.6)

Making use of the Equations 5.5 and 5.6, Equation 5.4 may be rewritten as

\[
\frac{\partial F}{\partial t} = \int_{v}^{\infty} K v^n g(v/v') \, dF(v',t)
\]  
(5.7)

A similarity variable \( z=(1+Kt)v^n \) exists for Equation 5.7 (Fillipov, 1961). For times sufficiently large this variable may be simplified to \( Ktv^n \). This implies that for a fixed value of \( F(v,t) \) (a horizontal line in Figures 7, 8 and 9) the similarity variable \( z=Ktv^n \) should remain constant if the above postulations are valid.
$Kv^nt=\text{constant}$ or $v^nt=\text{constant}$

Figures 13 and 14 are log-log plots of drop diameter $d$ vs time $t$ for dibenzyl ether and linseed oil systems at superficial gas velocities $u_g=0.1$ and 2 cm/sec respectively. It can be seen that the slopes of the straight lines are approximately $-0.33$. This means that the exponent $n$ in Equation 5.5 is approximately 1, because $v$ is proportional to $d^3$.

Using the approximate value $n=1$, cumulative distribution functions $F(v,t)$ are plotted against the similarity variable $t^{1/3}d$ for two different systems in Figures 15 and 16. It is shown that the cumulative distributions at various times collapse into a single line. The same behaviour was obtained for all three systems and experimental conditions in this work. Therefore, the existence of the similarity variable in drop breakage of lean dispersions in gas-liquid-liquid tubular reactors has been established. However, it has to be realized that the constant $K$ in Equation 5.5 may be a function of the local intensity of turbulence, operating and design parameters of tubular reactors. Hatzikiriakos et al. (1988c) found that $K$ may be related to the stable size spectrum, $d_{\text{max}}-d_{\text{min}}$, showing that $K$ is a function of the energy dissipation rate and physical properties of the dispersed phase liquid.

Knowledge of the similarity variable may lead someone in evaluating the density function. However, this is a numerical procedure and out of the scope of the present investigation. Details of the method for determining
the drop size distribution at various instants are given by Ramkrishan (1973).

5.3.3. Comparison with Agitated Vessels (Unsteady state)

Figure 17 is a plot of the ratio $d_{vs,t}/d_{vs}$ against time of column operation. The solid line represents Equation 5.2 and the dashed line represents the rate at which steady state is reached in agitated vessels with lean liquid-liquid dispersions (Narsimhan et al., 1980). The energy dissipation in the agitated vessel corresponds to superficial gas velocity of 4 cm/sec. As it may be concluded from Figure 17, steady state drop size distributions are approached more rapidly in tubular reactors than in agitated vessels.

Ramkrishna (1973) found that the transition probability function $\Gamma(v)$ in agitated vessels with lean dispersions is proportional to $d^6$. In Section 5.3.2 it was found that $\Gamma(v)$ in tubular reactors with lean liquid-liquid dispersions is proportional to $d^3$. The large exponent on the drop diameter in the case of agitated vessels indicates that the probability of drop break-up decreases more rapidly than in the case of tubular reactors as the drop size is reduced. This gives a reasonable explanation for why steady state is approached more rapidly in tubular reactors. It also suggests that tubular reactors perform better in the sense that larger interfacial areas are obtained more quickly. However, this is discussed in detail in Section
5.4. Maximum, Minimum and Mean Drop Sizes

Maximum, minimum and Sauter mean drop diameters are listed in Table 4 for the bulk of the experiments carried out in this investigation. Maximum and minimum stable drop sizes against break-up and coalescence set limits in the stability of drops in dispersion processes. As a result, a drop with size between these two limits is too small to break up and too large to coalesce. These sizes are also very useful in correlating the mean drop sizes. Sauter mean drop diameter is widely used in estimating the interfacial area. In this subsection the impact of superficial gas velocity and physical properties of dispersed liquid phase on the steady state Sauter mean diameter are discussed and generalized correlations for the maximum, minimum and mean drop sizes are presented. Also a new theory is proposed to account for the functional dependence of maximum and minimum drop sizes on the energy dissipation rates.

5.4.1. Superficial Gas Velocity

5.4.1.1. The Prediction of Mean Drop Size in Tubular Reactors for Gas-Liquid-Liquid Systems

Three flow regimes are clearly identified in the operation of bubble columns for gas-liquid-liquid systems as discussed in section 2.4. In the low gas flux regime, bubbles rise independently and the assumption that
local isotropic turbulence controls break-up is at best approximate. In addition the probability of direct bubble-drop collision is very low as drops tend to follow streamlines around bubbles as they move upwards through the column. The mechanism for drop break-up at low gas fluxes appears to be governed by shear stresses induced by bubble motion, where drop break-up arises from deformation and acceleration (Hatzikiriakos et al., 1988b). In APPENDIX B a simple expression is derived giving the maximum stable drop size against break-up as a function of superficial gas velocity in this flow regime:

\[
\sigma \quad \frac{d_{\text{max}}}{(\mu_c \rho g)^{0.5} \left( \frac{\mu_d}{\mu_c} \right) \mu g^{0.5}}
\]

The important part of this formula is that maximum drop diameter is expected to change as a function of -0.5 power of superficial gas velocity.

Following the Thomas approach (1980) in APPENDIX C, the minimum stable drop size against coalescence is found to be

\[
\frac{d_{\text{min}}}{(\sigma h/\mu_c)^{2/3} (v/g)^{1/6}}
\]

in the same flow regime. Again the minimum drop size against coalescence is a function of -0.5 power of superficial gas velocity. In APPENDIX D
arguments originating from a statistical point of view have shown that at low gas rates the superficial gas velocity is almost proportional to terminal velocity of bubbles.

In the high gas flux regime the flow field approaches local isotropic turbulent conditions. In this case Equations 2.9 and 2.10 for the maximum and minimum drop sizes against break-up and coalescence respectively may be applied. In APPENDIX A the energy dissipation in bubble columns has been found to be:

\[ \varepsilon = \frac{\Delta \rho g H u_g}{\rho_l H_o} \]  

(5.10)

where \( \Delta \rho \) is the density difference (continuous phase liquid-gas), \( \rho_l \) is the density of the liquid in the reactor, \( H_o \) is the initial liquid height and \( H \) is the overall pressure drop across the column. Equation 5.10 is based on the assumption that the energy is dissipated uniformly by the liquid in the reactor. The exact proportion absorbed by the dispersed phase cannot be calculated using this relation as a significant fraction of the energy dissipated is absorbed during drop break-up, through the creation of additional interfacial area. If all of the energy is transferred to the dispersed phase, then the power dissipated per unit mass of dispersed phase can be expressed as:
where \( \rho_{ld} \) is the density of the dispersed phase and \( \phi \) is the volume fraction of the dispersed phase. Experimentally one would anticipate that the power dissipated by the dispersed phase would fall between those obtained using Equations 5.10 and 5.11. Applying Equation 2.9, two maximum diameters, \( d_{\text{max},d} \) and \( d_{\text{min},u} \) can be obtained, where \( d_{\text{max},d} \) is the maximum drop size by assuming that the power is dissipated by the dispersed phase exclusively and \( d_{\text{max},u} \) is the maximum drop size by assuming that the power is dissipated uniformly by the liquid in the reactor. The maximum stable drop size against break-up would fall between the \( d_{\text{max},d} \) and \( d_{\text{max},u} \). Using similar arguments and Equation 2.10 two minimum diameters can be obtained, i.e., \( d_{\text{min},d} \) and \( d_{\text{min},u} \) (\( d_{\text{min},d} \) is the minimum stable drop diameter by assuming that the power is dissipated by the dispersed phase exclusively and \( d_{\text{min},u} \) is the minimum stable drop diameter by assuming that the power is dissipated uniformly by the liquid in the reactor). Mean drop diameter can be expressed as a function of minimum and maximum diameters obtained from Equations 2.9 and 2.10:

\[
d_{vs}=f(d_{\text{min},d}, d_{\text{min},u}, d_{\text{max},d}, d_{\text{max},u})
\]  

\[(5.12)\]
Postulating similar arguments discussed in Section 2.5 for the effect of relative viscosity $\mu_d/\mu_c$, Equation 5.12 may be rewritten as:

$$d_{vs} = f(d_{min,d}, d_{min,u}, d_{max,d}, d_{max,u})(\mu_d/\mu_c)^{0.25}$$

(5.13)

Equation 5.13 is used as a basis for correlating experimental drop size data in this study. The role of dispersed phase volume fraction is not given explicitly in Equation 5.13 since it has been incorporated in the calculation of minimum and maximum drop diameter, based on the assumption that the energy is dissipated by the dispersed phase exclusively.

5.4.1.2. Experimental results

Figure 18 shows the effect of superficial gas velocity on the steady state Sauter mean diameter of dispersed phase liquid drops for the systems used in this investigation. Two flow regimes, one below and one above 0.3 cm/sec are clearly identified by a sharp change in the slope of the functional dependence. For drop break-up to occur, it is necessary that enough energy be supplied to the drop to overcome the forces that resist breakage as a function of interfacial tension. The energy for the drop break-up may be kinetic energy at high gas fluxes (kinetic energy of energy-dissipating eddies), shear energy at low gas fluxes or a combination of the two. As was discussed in section 2.6, dynamic pressure fluctuations are the forces responsible for breakage of drops at high gas fluxes and
shear forces at low gas fluxes. One may expect that two correlations will be derived corresponding to high and low gas fluxes respectively. However, the practical importance of these processes lies at high gas fluxes (Hatzikiriakos et al., 1988a). In addition, most of the experiments carried out in this investigation are related to high superficial gas velocities. Due to these two reasons, an attempt was made to correlate one generalized equation to determine the mean drop size in terms of theoretical minimum and maximum drop sizes:

$$d_{vs} = 0.2 \left( d_{max,d} + d_{min,d} \right) + 0.04 \left( d_{max,u} + d_{min,u} \right)$$  \hspace{1cm} (5.14)

Equation 5.14 fits experimental mean drop diameter within ±30%. This deviation is slightly influenced by the incorporation of mean drop sizes at low gas fluxes. If this correlation for mean drop diameter is modified by including the ratio of dispersed phase viscosity to continuous phase viscosity, it becomes:

$$d_{vs} = (0.201(d_{max,d}+d_{min,d})+(0.0305(d_{max,u}+d_{min,u}))\left(\mu_d/\mu_c\right)^{0.256}$$  \hspace{1cm} (5.15)

Equation 5.15 correlates the Sauter mean diameter within ±22%. Close scrutiny of the size distribution data suggested that the effect of interfacial tension on mean drop diameter is overestimated by Equations 2.9 and 2.10 and an additional interfacial tension term is included in the correlation. Sauter mean drop diameter is predicted within ±10% with Equation 5.16:

$$d_{vs} = 0.15((d_{max,d}+d_{min,d})+0.7(d_{max,u}+d_{min,u}))\sigma^{-0.52}(\mu_d/\mu_c)^{0.256}$$  \hspace{1cm} (5.16)
where diameters are in microns and $\sigma$ is interfacial tension in dynes/cm. A parity plot for this correlation is shown in Figure 19. The values calculated from Equation 5.16 are very much comparable to those listed in Table 4.

From Figure 18 it may be seen that the cottonseed and linseed oil systems have identical behaviour and the two flow regimes can be identified clearly. However, the dibenzyl ether system with a relatively low viscosity in comparison with the other two systems under investigation, has the same behaviour at both low and high gas fluxes. This deviation may be attributed to a combined effect of relative viscosity and interfacial tension for systems of low relative viscosity and interfacial tension. Moreover, since the shear stresses are more important at low gas fluxes drop breakage rate depends on the interfacial tension and the shear field outside the drop. As the ratio $\mu_d/\mu_c$ is closer to unity the shear forces become more effective. Therefore, the Sauter mean diameters for such systems are smaller than expected (a slope -0.5 would give larger drop sizes).

Using the same arguments with those used for correlating the Sauter mean drop diameter, the experimental minimum diameter $d_2$ (diameter at 2% cumulative volume) and maximum diameter $d_{95}$ (diameter at 95% cumulative volume) were fitted within $\pm 10\%$ using Equations 5.17 and 5.18:

$$d_2 = 0.314(d_{\text{min},d} + 0.7d_{\text{min},u})\sigma^{-0.527}(\mu_d/\mu_c)^{0.256} \tag{5.17}$$
From Equations 5.16, 5.17 and 5.18 it can be observed that $d_{\text{max},d}$, $d_{\text{max},u}$, $d_{\text{min},d}$ and $d_{\text{min},u}$ appear with fixed proportionality constants. This suggests that the energy is dissipated in a fixed proportion by dispersed and continuous liquid phases. The magnitudes of proportionality constants imply that almost 60% of the available energy in the reactor is dissipated exclusively by the dispersed phase. This is understandable and due to the fact that most of the available energy is consumed to increase the interfacial area. However, the fixed percentage (60%) cannot be stated firmly due to the arbitrariness of some parameters e.g., the critical thickness for rupture was taken $h=10^{-5}$ cm, the proportionality constants in Equations 2.9 and 2.10 are not fixed (taking other constants a different percentage would be found). The coefficients of Equations 5.16-5.18 have been obtained through nonlinear least square regression of experimental data. The experimental maximum and minimum diameters have the same functional dependence on energy dissipation rate as mean drop diameter, as shown in Figures 20 and 21 for the cottonseed oil and linseed oil systems respectively. The various slopes related to this functional dependence for both flow regimes, are calculated using least squares and tabulated in Table 5. Therefore, the experimental maxima and minima may be correlated with high accuracy ($\pm 1\%$) as functions of Sauter mean diameters as follows:

\begin{equation}
    d_{\text{vs}} = 0.5 \; d_{g5} = 6 \; d_2 \tag{5.19}
\end{equation}
At gas fluxes below 0.3 cm/sec a slope of -0.5 can be noticed for both maximum and minimum diameters. This is in agreement with the slopes predicted by Equations 5.8 and 5.9. However at this point two questions related to the proposed theory have to be discussed before the theory is established. The first question is due to the consideration that drainage takes place between planes steadily pressed together for a period determined by the lifetime, \( T \), of maximum shear stress (the equator of a bubble where the stresses are concentrated, needs time equal to \( T = \frac{d_p}{U_T} \) in order to pass by a drop). One may wish to know, if the characteristic time for two colliding drops, \( t_o \), known as natural rebound time, is less than the characteristic time \( T \) defined in APPENDIX C for a two droplet encounter. If one presumed that \( T \) is always less than \( t_o \), then coalescence would always happen even if shear stresses are no longer configured. The characteristic time \( t_o \) is given by Lamb (1945) as:

\[
t_o = \left( \frac{\rho_g d_p^3}{\sigma} \right)^{1/2}
\]

(5.20)

Taking the ratio \( t_o/T \) using Equations 5.20 and \( T \sim \frac{d_p}{U_T} \),

\[
t_o/T = U_T \left( \frac{\rho_g d_p}{\sigma} \right)^{1/2}
\]

(5.21)

This ratio in all cases under investigation is less than unity, giving an answer to one's concern about the duration problem. For example, taking the surface tension to vary from 5 to 26 dynes/cm, \( d_p \) from 20 to 100 \( \mu \)m and
$U_T$ from 15 to 30 cm/sec, which includes all the cases under investigation, the ratio $t_o/T$ is always less than unity.

The second question arises from the following argument. Since the model assumes that the droplets are pressed together by a steady force and that the film is drained between plane solid surfaces, the predicted $d_{min}$ sizes should be smaller than the boundary layer thickness in the surroundings of the bubbles. For example, taking the size of the bubbles to vary from 1.3 to 2.2 cm (typical sizes encountered in the present work), the boundary layer thickness may be easily calculated to be in the range 230-260 μm. Experimental minimum drop sizes are in the range 25-90 μm. Additionally, by applying Equation C.6 for some typical values of $\sigma$, $\mu_c$, $h=10^{-5}$ cm and $U_T$ ($f(p)$ may be neglected, since in Taylor's expression changes a little for $5<p<19$ $f(p)=(19p+16)/(16p+16)$), minimum drop sizes are calculated to be in the range 25-65 μm. Therefore, these results justify the assumptions made for the present model. However, it has to be kept in mind that this theory is valid only for inviscid continuous phase and need to be modified for a more viscous continuous phase.

At high gas fluxes a slope of -0.25 is obtained for both maximum and minimum drop diameters. The -0.25 power law dependence of minimum drop diameter conforms with Equation 2.10 derived by Thomas (1981). However, one would expect a -0.4 power law dependence of maximum drop diameter on the rate of energy dissipation. This deviation may be attributed to the
following reasons:

- The Equations 2.9 and 2.10 have been derived under the assumption of isotropic turbulent conditions. The energy-dissipating eddies are responsible for break-up and coalescence of drops in isotropic turbulent flows. The length and velocity scales of these eddies are very much comparable with the minimum drop sizes. At these microscales, isotropic turbulent conditions may be reasonably considered. However, due to an order of magnitude difference in size between maximum and minimum drop diameters, isotropic turbulent conditions may not be considered at scales comparable to maximum drop sizes.

- The axial dispersion coefficient considerably exceeds the radial dispersion coefficient in tubular reactors (Reith et al., 1968). This finding is not consistent with conditions prevalent in isotropic turbulent fields.

- Zakrewski (1981) measured the intensity of turbulence in bubble columns and found that the spectral energy distribution follows a $k^{-2}$ law instead of $k^{-5/3}$ law predicted from Kolmogoroff theory, where $k$ is the wave number. In spite of this deviation from theory, Equations 5.17 and 5.18 predict the maximum and minimum drop sizes reasonably well by including the additional correction term of interfacial tension. This correlation derived under the consideration of isotropic turbulent conditions is very simple and appears to be applicable to a wide range of operating conditions.

Steady state drop size distribution as well as mean drop diameters are affected by changes in superficial gas velocity, as shown in Figures 22 and
23. The range of stable drop sizes narrows as the superficial gas velocity is increased. This has also been found by Shinnar (1961). As the energy dissipation rate increases and the maximum and minimum drop sizes become even more comparable in size, a dynamic equilibrium between coalescence and break-up occurs. A similar dependence is noted for agitated vessels (Stamatoudis and Tavlarides, 1985). The distributions of the number of drops over the drop size spectrum have a similar dependence on the energy dissipation rate as may be noticed from Figures 24 and 25. These distributions are not Gaussian but are severely skewed towards the smaller sizes. As the rate of energy dissipation increases, the number of smaller drops increases dramatically. The probability density becomes steeper in the smaller sizes due to the number of drops in this part of the drop size spectrum.

5.4.2. Viscosity

The impact of relative viscosity on the Sauter mean diameter is illustrated in Figure 26. The various systems used for sketching this Figure have almost the same interfacial tension (6.1-6.8 dynes/cm). Superficial gas velocity is a parameter. The slope of the curves are related to the power dependence of Sauter mean diameter on relative viscosity, \( \mu_d/\mu_c \). A power dependence of 0.256 was obtained by nonlinear regression on the experimental data which conforms with the value, 0.25, obtained by Calderbank (1967) for bubbles in agitated vessels. Such a dependence may
also apply to steady state Sauter mean diameter of bubbles in bubble columns under certain operating conditions (Shah et al., 1982). Nishikawa et al. (1987) reported a power dependence ranging from 0.125 to 0.2 for liquid-liquid systems in agitated vessels.

As it is seen from Figure 26, the dependence of the steady state Sauter mean diameter seems to be the same at both low and high gas fluxes. The existence of different mechanisms of drop splitting at different flow regimes does not appear to affect this dependence. Using least squares on each line appearing on Figure 26, various slopes were calculated with a standard deviation of 5%. At low gas fluxes where shear forces dominate, the ratio of drop viscosity to that of the suspending fluid is an important dimensionless group in governing drop deformation and burst (Rallison, 1984). This group plays the role of matching the normal and tangential stresses at the liquid-liquid interface. As this ratio deviates from the range of 0.1 to 1 to either higher or lower values, drop resistance to break-up becomes greater and greater. These theoretical arguments conform with studies on dispersion phenomena at simple flows like couette and hyperbolic flows (Grace, 1982) and at those of the present work. At high gas fluxes where dynamic pressure fluctuations dominate, the dependence of the steady state Sauter mean diameter on relative viscosity is also consistent with theory (Hinze, 1955). There is no experimental evidence related to the dependence of Sauter mean diameter on relative viscosity for gas-liquid-liquid systems in tubular reactors.
It has to be pointed out that at high gas fluxes and for the dibenzyl ether system (low interfacial tension and viscosity) a milky emulsion is formed. Photographs could not be taken. However, an expected Sauter mean diameter is close to 100 µm, as may be seen from the extrapolated line in Figure 26.

The role of viscosity is not limited to the ratio of drop viscosity to that of the suspending fluid. The viscosity of continuous phase itself is also of great importance. The time \( \tau \) required for the film of a two-drop encounter to drain to the critical rupture thickness \( h \) decreases as the viscosity of continuous phase decreases. In such a case, the rate of drop coalescence increases, as a result, bigger drops may be observed (Stamatoudis and Tavlarides, 1985).

5.4.3. Interfacial Tension

Interfacial tension has a limited influence on mean drop diameter as shown in Figure 18. The systems cottonseed oil and linseed oil, for example, have approximately the same viscosity but different interfacial tensions. The effect of interfacial tension on mean drop size is less than anticipated by the Equations of Thomas and Hinze which are based on isotropic turbulence. This difference may be attributed to the efficiency and modes of energy transfer to the dispersed phase which may not conform with conditions prevalent in isotropically turbulent flow fields. This was discussed in Section 5.4.1.b and it was concluded that isotropic turbulent conditions are not consistent with conditions prevalent in bubble columns.
at least in length scales comparable to maximum stable drop sizes against break-up. Equation 5.15 based on isotropic turbulence conditions fits the experimental Sauter mean drop diameters within ±22%. Once the additional term of interfacial tension $\sigma$ is included, Equation 5.16 predicts Sauter mean diameter within ±10%.

From Table 4, it may be seen that the effect of interfacial tension on Sauter mean diameter as well as on maximum and minimum drop diameters is stronger at low gas fluxes than at high gas fluxes. The differences in $d_{vs}$, $d_{max}$, and $d_{min}$ for the systems cottonseed oil and linseed oil, for example are comparatively larger at low gas fluxes. This is understandable if one takes into consideration that at low gas fluxes the interfacial tension forces oppose the shear forces in deforming the drops. Therefore a conflict between shear forces and interfacial tension forces exist and the drop breakage is expected to depend on the surface tension and on the hydrodynamic field outside the drop. At high gas fluxes, one has to consider the problem from an energy point of view. The deformation and bursting of drops are continuous processes since the drops are under the action of dynamic pressure fluctuations at every instant. Therefore, drops may promote larger deformations. For highly deformed drops, interfacial tension is in favor with splitting instead of resisting to it (Rallison, 1984). Since there is no limitation in energy, the assumption of the existence of highly deformed drops in turbulent fields is reasonable. The impact of interfacial
tension is also discussed below in section 5.5, where the tubular reactors are compared with the agitated vessels.

5.4.4. Dispersed Phase Concentration

The effect of dispersed phase concentration on Sauter mean diameter is shown in Figures 27 and 28 for linseed oil and cottonseed oil systems respectively. The solid lines on these Figures represent Equation 5.16. It may be seen that this correlation can predict the effect of dispersed phase concentration very well. It is also shown that Sauter mean diameter increases with the dispersed phase concentration. This dependence is expected to be stronger for denser dispersions (Φ > 0.10) since the coalescence rate of drops increases with concentration (Mlynek and Resnick, 1972; Coulaloglou and Tavlarides, 1976). However, for lean dispersions (Φ < 0.0045), it was found that the dispersed phase concentration had no effect on Sauter mean diameter. A similar behavior is noted for agitated vessels (Narsimhan et al., 1976). A limited dependence on concentration has also been reported by Nishikawa et al. (1987) for dense liquid-liquid dispersion in agitated vessels. Kato et al. (1984) reported that at Φ > 0.05 concentration had little impact on Sauter mean diameter in gas-liquid-liquid bubble columns.

From Figures 27 and 28, it may also be inferred that the impact of volume fraction of dispersed phase on Sauter mean diameter is the same at both low and high gas fluxes. The role of dispersed phase concentration is
not given explicitly in Equation 5.16 since it has been incorporated in the calculation of minimum and maximum drop diameters based on the assumption that the energy is dissipated by the dispersed phase exclusively.

5.4.5. Density

The relative density of the dispersed and continuous phases, $\rho_{ld}/\rho_{lc}$, had no effect on Sauter mean diameter in the range $0.95 < \rho_{ld}/\rho_{lc} < 1.05$. In Table 6, the Sauter mean diameters for three subsets of experiments are listed. In each subset the superficial gas velocity and interfacial tension are kept constant. The differences in $d_{vs}$ may be attributed to relative viscosity which has a significant effect on $d_{vs}$ as it is discussed in Section 5.4.2. Therefore, the Sauter mean diameters have to be corrected using Equation 5.16 in order to account for the different relative viscosities. In Table 6, the corrected Sauter mean diameters are listed in contrast with the real Sauter mean diameters. Basis is taken as $\mu_d/\mu_c = 13$, an intermediate value. As it may be observed, the differences for the first subset of experiments are very small and these can be attributed to experimental error or the effect of interfacial tension. However, the data for the other two subsets of experiments is inconclusive.

Hinze (1955) concluding his classical paper pointed out that the difference in density between the dispersed and the continuous phases has an important effect on the way in which break-up occurs. Thus, it has to be
realized that correlations for Sauter mean diameters using the maximum and minimum drop diameters should be based on data without a basic difference in densities between the two phases. If there is, incorrect results might be obtained. This turns out not to be the case, only if the role of relative density has to be assessed independently. Following these concluding remarks of Hinze, data with difference in the density of the two liquid phases are not included in the correlations 5.16-5.18.

5.4.6. Other Parameters

The initial drop size has no effect on the steady state Sauter mean diameter for initial drop sizes in the range 2000μm<d<10000μm. However, the initial drop size have an impact on the drop breakage rate for times less than about 3 min, as it is discussed in Section 5.3.1.

Operating the column under constant pressure, constant gas rate or intermediate conditions, different bubble sizes are obtained at constant gas fluxes. It was found that bubble sizes in the range 10 to 25 mm had no apparent effect on mean drop diameter. However, it has to be emphasized that at high gas fluxes, the photographic technique used in this investigation is limited to the bubbles in the vicinity of the column wall. At these high gas fluxes, the large bubbles tend to rise in the central plume of the column. Therefore, the upper limit of 25 mm is a little higher in reality.

A perforated plate was placed 5 cm above the nozzles in order to avoid
drop break-up because of the air jet, in particular at high gas fluxes. However, such an effect was not observed and the rest of the experiments were performed following the procedure described in Section 3.2.

Finally, a perforated plate was placed 5 cm below the free surface of continuous liquid phase in order to avoid free surface effects on drop break-up, which might have an impact on the Sauter mean diameter.

5.5. Comparison with Agitated Vessels (Steady State)

Agitated vessels are widely used in Chemical Engineering as contactors for liquid-liquid systems because of their effectiveness in bringing about the contact of two liquid phases. Their characteristics are discussed in Section 2.5. On the other hand, bubble columns are also used extensively for gas-liquid systems because of low investment costs and effective mixing of fluids. Their characteristics are presented in Section 2.3.

In this section an attempt is made to compare the relative performance of tubular reactors and agitated vessels as contactors for liquid-liquid systems. This comparison is based on the energy consumption per unit of interfacial area. In particular, the results of this investigation are compared with those of Nishikawa et al. (1987), who summarized the impact of operating parameters and physical properties of both liquid phases on the Sauter mean diameter of dispersed phase drops in agitated vessels operating at steady state with Equations 2.13 and 2.14 given in section 2.5.

The impact of power dissipation on the Sauter mean diameter for
agitated vessels and tubular reactors is illustrated for three different systems in Figure 29. For linseed oil system, Figure 29.a, a system with high viscosity and low interfacial area, the mean diameters $d_{vs.a}$ and $d_{vs.b}$ obtained in an agitated vessel and a tubular reactor respectively are comparable. The ratio $d_{vs.a}/d_{vs.b}$ converges at high energy dissipation rates. The large mean diameters obtained at low energy dissipation rates in agitated vessels can be attributed to coalescence in zones remote from the impeller (Mlynek and Resnick, 1972; Weinstein and Treybal, 1973).

Figure 29.b shows the effect of energy dissipation rate on Sauter mean diameter for the cottonseed oil system, a system with high viscosity and high interfacial tension. It can be observed that mean drop diameter is much greater in agitated vessels as far as liquid-liquid systems are concerned. This may be attributed to the local energy dissipation and accordingly to the intensity of turbulence which is more uniform in the tubular reactors. In the agitated vessels, the Sauter mean diameter might be affected significantly by the local intensity of turbulence in the vicinity of the impeller. The zones remote from the impeller in the tank remain less effective for liquid-liquid systems. To gain a better understanding of this fact, it has to be realized that the drop deformation and breakage are time-dependent processes. In agitated vessels the drops circulate from zones of high energy dissipation to zones of low energy dissipation. Therefore, there may not be enough time in order for the drops to be
deformed critically. On the other hand in tubular reactors the drops are under the action of dynamic pressure fluctuations for longer times. Therefore, the required energy for drop bursting is accumulated easier. Yoshida and Yamada (1972) performed experiments in both tubular reactors and agitated vessels for dispersion of kerozene in water. This system has high interfacial tension (43 dynes/cm) and low viscosity (1.29 cp). It was found that the Sauter mean diameter of kerozene droplets in the tubular reactor was approximately one half of that in the agitated vessel. These findings are consistent with those of the present work.

Figure 29.c shows the same comparison for the dibenzyl ether system, a system with low viscosity and interfacial tension. In this case, agitated vessels and tubular reactors perform equally well. Clearly, the comparative advantage for tubular reactors is greatest for fluids with high interfacial tension as the steady state Sauter mean diameter has only a limited dependence on interfacial tension in these reactors.
6.0. CONCLUSIONS

A number of characteristics related to gas-liquid-liquid systems in tubular reactors were investigated. Tubular reactors were compared with liquid-liquid agitated vessels on the basis of their relative performance as phase-contacting equipment. The results of this investigation may be summarized as:

1. **Gas Flux:** as it was expected the gas flux plays an important role in determining the steady state Sauter mean diameter of dispersed phase drops. It was found that Sauter mean diameter is decreased more significantly with gas flux at low gas rates. Two flow regimes, one below and one above ~ 0.3 cm/sec, were identified on a plot $d_{ve}$ vs $\varepsilon$ by a sharp change in the slope of the functional dependence.

2. **Physical Properties of Dispersed Phase Liquid:** the physical properties of dispersed phase constituents have a significant impact on the steady state drop size and drop size distribution. Generally steady state Sauter mean diameter increases strongly with relative viscosity, moderately with dispersed phase volume fraction and slightly with interfacial tension. The relative density was not found to have an effect in the range 0.95 to 1.05 g.cm$^{-3}$. The steady state Sauter mean diameter was correlated as a function of gas flux and physical properties of dispersed phase liquid. This correlation predicts mean drop sizes within ±10%.

3. **Maximum and Minimum Stable Drop Sizes:** a new theory was proposed to
explain the functional dependence of maximum and minimum stable drop sizes against break-up and coalescence respectively at low gas rates on the energy dissipation rate. Two formulae were derived corresponding to maximum and minimum stable drop sizes which give an approximation of the order of magnitude of these sizes at low gas fluxes. At high gas fluxes the functional dependence was attempted to be explained with the isotropic turbulence theory of Kolmogoroff. However, it was found that bubble columns are not quite isotropic even at microscales comparable to maximum drop sizes (500 μm).

4. Unsteady State Drop Size Distribution: the rate at which steady state is reached was found to be independent of the superficial gas velocity and the physical properties of the dispersed phase liquid. The reduction in mean drop size was correlated as a function of time. The existence of a similarity variable was identified and suggestions were made for modeling the unsteady state drop size distribution as a function of time. The transition probability function was found to be proportional to drop volume $v$ in contrast with $v^2$ in the agitated vessels.

5. Comparison with Agitated Vessels: as mean drop diameter is less dependent on interfacial tension in tubular reactors than in agitated vessels, the comparative advantage is greatest for two phase liquids with high interfacial tensions. Tubular reactors are also preferred for two phase liquids with a high relative viscosity, at low energy dissipation rates. A comparison between tubular reactors and agitated vessels operating under
unsteady state conditions showed that tubular reactors can perform better for all liquid-liquid systems. This is attributed to different functional dependence of the transition probability function on the droplet volume in tubular reactors and agitated vessels.
7.0. RECOMMENDATIONS

The results of this investigation suggest that studies to the following areas are warranted:

1. The isotropic turbulent theory of Kolmogoroff generally gives simple models, applicable to a wide range of operating conditions. However, this theory is sometimes hidden by empiricism. The efficiency and modes of energy dissipation have to be investigated further, particularly in tubular reactors where it was found that these reactors are not quite isotropic even at scales comparable to maximum drop sizes (500 \( \mu \text{m} \)).

2. The continuous phase viscosity \( \mu_c \) itself is expected to have a strong impact on the steady state mean drop size. As \( \mu_c \) increases, bubbles slow down and the required time \( \tau \) for the film in a two-drop encounter to drain to the rupture thickness \( h \) increases. Therefore, coalescence rates will be important only at high volume fractions of dispersed phase.

3. In lean liquid-liquid dispersions, the effect of drop coalescence is negligible. Therefore, to model the transient droplet breakage only the effect of break-up has to be considered. However, in dense liquid-liquid dispersions population balances have to include the effect of coalescence as well. Dynamic relationships between break-up and coalescence have to be found and checked with experimental data.

4. Interfacial area has a direct impact on mass transfer processes. Gas-liquid and liquid-liquid interfacial areas are expected to play a role for
mass transfer from the gas phase to dispersed liquid phase. The mechanism or the mechanisms of mass transfer in G-L-L systems should be studied.

5. Gas-bubble plate interactions in multi-stage tubular reactors are expected to improve mass transfer rates and impose local axial mixing patterns for the liquid phase above and below the plate. However, the existence of plates may impose milder conditions than those without plates as far as the gas flux remains the same. This may lead to a decrease in interfacial area or an increase in gas requirements. All these effects have to be quantified in terms of process variables in order to be applied to the design of a novel tubular reactor.
8.0. NOTATION

A  cross sectional area of tubular reactor
Bo  bubble Bond number, $=\frac{\rho g \alpha^2}{\sigma}$
C  proportionality constant in Equation 2.11
d  diameter of impeller in Equations 2.13 and 2.14
dp  drop diameter, $\mu m$
dv  Sauter mean drop/bubble diameter, $\mu m$
dva  Sauter mean diameter (agitated vessel), $\mu m$
dvb  Sauter mean diameter (tubular reactor), $\mu m$
dv,t  Sauter mean diameter at time t, $\mu m$
dmin  minimum stable drop size, $\mu m$
dmax  maximum stable drop size, $\mu m$
dmin,u  minimum drop diameter (uniform energy dissipation), $\mu m$
dmin,d  minimum drop diameter (assuming all energy is dissipated to the dispersed phase), $\mu m$
dmax,u  maximum drop diameter (uniform energy dissipation), $\mu m$
dmax,d  maximum drop diameter (assuming all energy is dissipated to the dispersed phase), $\mu m$
d2  experimental minimum diameter at 2% cumulative vol.%, $\mu m$
d95  experimental maximum diameter at 95% cumulative vol.%, $\mu m$
db  diameter of a spherical bubble, cm
d  equivalent diameter of a bubble, cm
D  diameter of impeller in agitated vessels, cm; diameter of agitated vessel in Equations 2.13 and 2.14, cm
Do  diameter of the orifice, cm
f(p)  function of relative viscosity
F(v,t)  volume fraction cumulative distribution function of drop size
Fr  bubble Froude number, $Fr=U_T\sqrt{\frac{\rho}{\Delta \rho gd_b}}$
F  force, dynes
g  gravitational acceleration constant, $\text{cm.s}^{-2}$
G  shear rate, $\text{s}^{-1}$
G(v,v')  cumulative distribution function of daughter droplet size from break-up of a large drop
h  critical rupture thickness, $\approx 10^{-5}$ cm
H  liquid depth, cm
\( H_o \)  
 initial liquid depth, cm

\( k \)  
 wave number, \( k = 2\pi/\lambda \), cm\(^{-1}\)

\( K \)  
 constant in Equation 5.6

\( n \)  
 constant in Equation 5.6

\( N \)  
 number of revolutions of impeller per unit time in Equation 2.11; number of bubbles rising per unit time

\( N_{c} \)  
 dimensionless capacitance number

\( N_{w} \)  
 dimensionless number, \( N_{w} = Bo Fr \cdot 5 \)

\( p \)  
 relative viscosity of continuous liquid phase to dispersed liquid phase, \( p = \mu_d/\mu_c \)

\( P \)  
 pressure, cm of continuous liquid phase

\( P_{h} \)  
 hydrostatic pressure at orifice plate, cm of continuous liquid phase

\( Re \)  
 bubble Reynolss number, \( Re = U_T d_e \rho_f/\mu_c \)

\( s \)  
 film radius in a two-drop encounter, cm

\( t \)  
 time, min

\( t_o \)  
 natural rebound time of a drop, s

\( T \)  
 characteristic time scale, s

\( u \)  
 velocity scale of energy-dissipating eddies, cm.s

\( u_s \)  
 slip velocity of a bubble swarm, cm.s

\( u_g \)  
 superficial gas velocity, cm.s

\( U_T \)  
 terminal velocity of bubble, cm.s

\( v \)  
 drop volume, cm\(^3\)

\( v' \)  
 drop volume, cm\(^3\)

\( V_c \)  
 volume of the chamber, cm\(^3\)

\( z \)  
 similarity variable, \( = K t v^n \)

**GREEK LETTERS**

\( \alpha \)  
 equivalent radius of bubble, cm

\( \Gamma(v) \)  
 transition probability function characterizing droplet breakage

\( \Delta p \)  
 difference in density between continuous phase liquid and gas, g.cm\(^{-3}\)

\( \varepsilon \)  
 energy dissipation per unit mass, cm\(^2.s^{-3}\)

\( \varepsilon_d \)  
 energy dissipation per unit mass of dispersed phase liquid, cm\(^2.s^{-3}\)

\( \varepsilon_o \)  
 critical energy dissipation per unit mass, cm\(^2.s^{-3}\)

\( \varepsilon_g \)  
 gas holdup
<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\eta$</td>
<td>length scale of energy dissipating eddies, cm</td>
</tr>
<tr>
<td>$\theta_w$</td>
<td>the wake angle behind a bubble</td>
</tr>
<tr>
<td>$\lambda$</td>
<td>wavelength, cm</td>
</tr>
<tr>
<td>$\mu_c$</td>
<td>viscosity of continuous phase, g.cm$^{-1}$.s$^{-1}$</td>
</tr>
<tr>
<td>$\mu_d$</td>
<td>viscosity of dispersed phase, g.cm$^{-1}$.s$^{-1}$</td>
</tr>
<tr>
<td>$\nu$</td>
<td>kinematic viscosity of continuous phase liquid, cm$^2$.s$^{-1}$</td>
</tr>
<tr>
<td>$\rho_l$</td>
<td>density of the continuous phase liquid, g.cm$^{-3}$</td>
</tr>
<tr>
<td>$\rho_d$</td>
<td>density of dispersed phase liquid, g.cm$^{-3}$</td>
</tr>
<tr>
<td>$\rho_g$</td>
<td>density of gas, g.cm$^{-3}$</td>
</tr>
<tr>
<td>$\sigma$</td>
<td>interfacial tension, dynes/cm</td>
</tr>
<tr>
<td>$\sigma_{g-l}$</td>
<td>gas-liquid surface tension, dynes/cm</td>
</tr>
<tr>
<td>$\tau$</td>
<td>time required for the film to drain to thickness $h$, s</td>
</tr>
<tr>
<td>$\phi$</td>
<td>volume fraction of dispersed phase liquid</td>
</tr>
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9.0. REFERENCES


APPENDIX A

Calculation of the Energy Dissipation Rate in Tubular Reactors

The energy balance between the bottom and the top of the reactor can be used to determine the energy dissipation rate. The energy at the bottom includes the kinetic, potential and surface energy of bubbles. The energy at the top includes the kinetic and surface energy of bubbles. If it is assumed that the bubbles do not break up or coalesce while passing through the reactor, then the change in surface energy can be taken as zero. The velocity of bubbles also does not change significantly during the passage through the reactor, this indicates that the change in kinetic energy is negligible. Hence, the energy dissipated in the reactor with an initial height of $H_0$ cm is as follows:

$$\Delta E = E_{\text{bottom}} - E_{\text{top}}$$

where

$$E_{\text{bottom}} = \frac{1}{2} m u_1^2 + \Delta \rho g H Q,$$

$$E_{\text{top}} = \frac{1}{2} m u_1^2,$$

$m$ is the mass flow rate of gas and $Q$ is the volumetric flow rate of gas.

Taking into account the above stated assumptions,

$$\Delta E = \Delta \rho g H Q$$

The energy dissipation rate per unit mass of liquid becomes:
\[
e = \frac{\Delta \rho \, g \, H \, u_g}{\rho_1 \, H_o}
\]  
(A.1)

Where \(\Delta \rho\) is the density difference between the continuous phase liquid and gas, \(\rho_1\) is the density of the continuous phase liquid and \(H\) is the head pressure in cm of liquid in the reactor.
APPENDIX B

Maximum Drop Size against Break-up at Low Gas Fluxes

When a bubble is rising independently in a low viscosity liquid, a boundary layer is formed around its surface. The equations governing the boundary layer have been studied by Moore (1962, 1965). It has been shown that the thickness of the boundary layer is proportional to $d_e Re^{-0.5}$, where $d_e$ is the equivalent diameter of the bubble and $Re$ is the Reynolds number. It has also been shown that the velocity changes within the order of magnitude of the terminal velocity of the bubble. In a flow field where bubbles are rising independently, drops follow the streamlines. Head-on collisions are not excluded but the probability of such events is very small. Moore (1965) also argued that the shear stresses are concentrated around the equator of bubbles. A drop in such a shear field tends to rotate, accelerate and deform. If the velocity gradients are large enough and interfacial tension forces are no longer able to maintain the drops intact they rupture. Taking into consideration all the above statements, the shear rate is of the order:

$$G \sim \frac{U_T}{d_e Re^{-0.5}}$$  \hspace{1cm} (B.1)

Substituting $Re = U_T d_e \rho / \mu_c$ into Equation B.1 and combining it with the Equation 2.2, Equation B.1 may be rewritten as:
Taylor (1934) argued that a drop of size $d_p$ in a simple shear field will break up if the ratio of viscous forces over interfacial tension forces exceeds unity. Therefore, a criterion for break-up is:

$$G \mu_c f(p) > \sigma/d_p$$

where $G$ is the shear rate taken from Equation B.2, $f(p)$ a function accounting for the relative viscosity and $p$ is the relative viscosity, $\mu_d/\mu_c$. After some manipulation Equation B.3 may be expressed as:

$$d_{\text{max}} \sim \frac{\sigma}{(\mu_c \rho_f g)^{0.5} f(p) U_T^{0.5}}$$

At low gas fluxes $U_T \propto u_g$ (see APPENDIX D). Therefore, Equation B.4 can be rewritten as:

$$d_{\text{max}} \propto \frac{\sigma}{(\mu_c \rho_f g)^{0.5} f(p) u_g^{0.5}}$$
Equation B.4 gives the order of magnitude of maximum stable drop size against break-up in tubular reactors at low gas fluxes. Equation B.5 includes the qualitative information that $d_{\text{max}}$ is expected to change as a function of $u_g^{-0.5}$. 
APPENDIX C

Minimum Drop Size against Coalescence at Low Gas Fluxes

The mechanism of coalescence includes the two following stages:
- The draining of the intervening film of continuous phase liquid to a critical thickness $h$.
- The rupture of the remaining film by a mechanism not completely understood.

It will be supposed that the two drops are pressed together by a steady force $F$, which is related to viscous forces and that drainage takes place between rigid planes. Calculation of the time $\tau$ required for the film to drain to the thickness $h$ is a classical lubrication problem and the solution is given by Thomas (1981) as:

$$\tau = \frac{3\pi \mu_c s^4}{2F h^2} \quad (C.1)$$

where $s$ is the radius of the film, $\mu_c$ the viscosity of the continuous phase liquid and $F$ is the force pressing the drops together. Considering the pressure within the drop gives:

$$F = 4\pi s^2 \sigma / d_p \quad (C.2)$$

so the film-drainage time may be rewritten as:
The viscous forces may be expressed as:

\[ F = \pi d_p^2 \mu_c G_f(p) \]  
(C.4)

Combining Equations C.2, C.3 and C.4, Equation C.4 can be expressed as:

\[ \frac{3}{32\pi} \mu_c \pi d_p^2 G_f(p) \left( \frac{d}{\sigma h} \right)^2 \]  
(C.5)

If one denotes by \( T \) the characteristic timescale of a two-drop encounter and assuming that viscous forces are the cause of drop movement, dimensional analysis yields:

\[ T = \frac{d_p}{U_T} \]

where \( d_p \) is the diameter of drop and \( U_T \) is the terminal velocity of bubble. A simple criterion for coalescence is \( \tau < T \), which gives after some manipulation:

\[ d_{\text{man}} \sim 2.2 U_T^{-0.5} f(p)^{-1/3} (\sigma h/\mu_c)^{2/3} (v/g)^{1/6} \]  
(C.6)

At low gas fluxes \( U_T \sim u_g \) (see APPENDIX D). Therefore, Equation C.6 may be rewritten as:

\[ d_{\text{min}} \propto u_g^{-0.5} f(p)^{-1/3} (\sigma h/\mu_c)^{2/3} (v/g)^{1/6} \]  
(C.7)
Equation C.6 gives the order of magnitude of minimum stable drop size against coalescence in tubular reactors at low gas fluxes. Equation C.7 includes the qualitative information that drop size is expected to change as a function of $u_g^{-0.5}$. 
APPENDIX D

Relationship Between Superficial Gas Velocity and Terminal Velocity of Bubbles at Low Gas Fluxes

Suppose that at an instant in the tubular reactor the superficial gas velocity is $u_g$, the number of bubbles per unit of time $N$ and that the bubbles are of uniform diameter $d_b$ and volume $V_b$. If the superficial velocity increases to $u_g + du_g$, where $du_g$ represents an infinitesimal increment, then the number of bubbles per unit of time will be $N + dN$ of diameter $d_b + dd_b$ and volume $V_b + dV_b$.

Writing down a mass balance for the gas and considering constant density, one gets:

$$Au_g = NdV_b + dN(V_b + dV_b)$$  \hspace{1cm} (D.1)

where $A$ is the cross-section area. Cancelling out the term $dNdv_b$ and integrating using $u_g = 0$ when $N = 0$, Equation D.1 may be rewritten as:

$$u_g = \frac{NV_b}{A}$$  \hspace{1cm} (D.2)

Now if one considers a swarm of bubbles covering completely the volume of the reactor, then the probability including both events for a head-on collision between a drop and bubble and a drop being trapped in the boundary layer of a bubble, is unity.

$$p = 1 \quad \text{and} \quad N' = V/V_b$$  \hspace{1cm} (D.3)

where $p$ represents probability, $N'$ number of bubbles and $V$ the volume of the
tubular reactor. When \( N \) bubbles are present in the reactor, then the probability becomes:

\[
p = \frac{N}{N} \quad \text{or} \quad p = \frac{NV_b}{V}
\]  \hspace{1cm} (D.4)

If the superficial gas velocity \( u_g \) increases, the number of bubbles will increase to \( N + dN \) and their volume to \( V_b + dV_b \). Then the probability will be:

\[
p + dp = \frac{(N + dN)}{N} \]

Using Equations D.3, D.4 and D.5 and integrating using that \( p = 0 \) when \( N = 0 \), one gets:

\[
p = \frac{NV_b}{V}
\]  \hspace{1cm} (D.6)

Combining Equations D.5 and D.6 one ends up with:

\[
p \propto u_g
\]  \hspace{1cm} (D.7)

Now consider that the number of drops per unit volume of the reactor is \( \eta \) and a bubble moving independently with its terminal velocity \( U_T \). Within a fixed time \( t \) this bubble will travel a distance \( l_1 = U_T t \) and have a probability \( p \) to collide with a drop or to entrap a drop in its boundary layer. For another bubble moving with velocity \( U_T + dU_T \), the covered distance during the same time will be \( l_2 = (U_T + dU_T)t \) and the probability \( p + dp \). If one considers a uniform distribution of drops in the reactor, then:

\[
l_1/l_2 = \eta_1/\eta_2 = p/(p + dp) = U_T / (U_T + dU_T)
\]  \hspace{1cm} (D.8)

Integrating Equation D.8, one may write:
\[ p \propto U_T \]  \hspace{1cm} \text{(D.9)}

Combining Equations D.7 and D.9, the following relationship can be obtained:

\[ u_g \propto U_T \]  \hspace{1cm} \text{(D.10)}

From a statistical point of view, Equation D.10 holds reasonably well. Sriram (1977) has defined the superficial gas velocity given in section 2.3 as Equation 2.7. If consideration of uniform bubble size is employed, then Equation D.11 may be obtained:

\[ u_g = \epsilon_g U_T \]  \hspace{1cm} \text{(D.11)}

Since gas hold-up does not change much and may be considered almost constant for superficial velocities in the range (0.1-0.3 cm/sec), the superficial gas velocity \( u_g \) can be considered as proportional to terminal velocity of uniform size bubble \( U_T \).