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A STUDY OF METHODS FOR PREDICTING THE CHANGE IN AZEOTROPIC COMPOSITION WITH PRESSURE AND WITH TEMPERATURE

BY

ADOLF PRIBUSH

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INTRODUCTION

Much data can be found in the literature giving the pressure-temperature-composition relationship for azeotropic systems at one pressure, usually atmospheric. Comparatively few data, however, have been published giving compositions over a range of pressures and temperatures for azeotropic systems. Such data may be useful in application to the separation of liquid mixtures by azeotropic distillations, or even simple fractionations, at pressures other than atmospheric. It may be found, for instance, that a particular system that forms an azeotrope at atmospheric pressure (and therefore cannot be separated beyond the azeotropic composition by means of a simple fractionation at atmospheric pressure, enabling a complete separation by means of a simple fractionation.

In the absence of complete pressure-temperaturecomposition data for many azeotropic systems, it is
desirable to have reliable mathematical relationships
available for predicting such data. A number of
relationships for predicting the change in azeotropic
composition with change in pressure and temperature
can be found in literature, but not all of them have
been checked sufficiently against experimental data to
enable their use with a knowledge of the degree of

accuracy that can be expected.

Object

The object of this paper is to present the results of a study of methods for predicting the change in azeotropic composition with pressure and with temperature, and to indicate the usefulness of the methods.

Scope

A study was made of the types and formation of azeotropes, the general effect of pressure and temperature on azeotropes, and published methods for predicting the effect of pressure and temperature on the composition of azeotropes. Each of these phases of study is presented in consecutive order to supply background information.

perimental data. Each method was tested with experimental data for ten assotropic systems for which the most complete data could be found in literature. Only binary systems were used. The systems included maximum and minimum-boiling homogeneous assotropes, and one minimum-boiling heterogeneous assotrope. One of the methods tested represents a pressure-composition relationship, and the other represents a temperature-composition relationship.

The information presented in this paper is based on

a literature search covering the years 1987-1953.

Materials and Equipment

All calculations were made with a calculating machine in conjunction with a five-place logarithm table. The calculations are reported to at least one place beyond that of the data found in the literature. Where vapor pressure data for pure components appeared with azeotropic data, it was used in the calculations made for its respective azeotropic system for the sake of consistency. Where vapor pressure data for pure components did not appear with their respective azeotropic systems, published vapor pressure tables were used.

Procedures

In the case of the equation tested for predicting the azeotropic temperature-composition relationship, the constants were determined by two methods. The method of least squares was used for those systems having ten or fewer points, and the method of averages was used for those systems having more than ten points.

The vapor pressure data for pure components used in the equation tested for predicting assotropic pressure-composition relationship was obtained from the sources mentioned above, using a log vapor pressure versus 1/(°C/230)

relationship.

Findings

rhe findings, and the azestropic and pure component vapor pressure data obtained from the literature, are presented in table form for easy reference.

Plan of Thesis

formation concerning definitions, theory of formation, and relationship of pressure, temperature, and composition, of azeotropes for background information. A short review is then given of published methods for predicting the change in composition of azeotropes with the change in pressure and temperature, indicating the nature of their derivation, references, and expected accuracy. The results of testing two equations for adequacy in predicting the change in composition of azeotropes with the change in pressure and temperature are then presented and evaluated. In the final section of the text, conclusions are given concerning the work covered by this paper.

DEFINITIONS AND THECKY OF AZEOTROPE FORMATION Definitions

An azectrope is characterized by a constant boiling point and composition, at a fixed pressure, of a mixture of two or more liquid compounds. It is also called a constant boiling mixture (C.B.M.). An azectrope cannot be separated into its components by a simple fractionation at a given pressure, since the composition and boiling point will remain constant.

some azertropes exist as one liquid phase (homogeneous mixtures) and others exist as two or more liquid phases (heterogeneous mixtures). This serves as a basis for the classification of azertropes. They are further subclassified as minimum-boiling or positive azertropes, and maximum-boiling or negative azertropes. The positive and negative classifications indicate that the mixture shows positive, or negative, deviations from ascult's law. Homogeneous azertropes have been found to exist as minimum and maximum-boiling mixtures. All reported heterogeneous azertropes are minimum-boiling mixtures (16).

Pormation of Azeotropes

The partial pressure of a component in equilibrium with a solution in a binary system that adheres to Haoult's law will be directly proportional to its mole fraction in the system. For such a system, the vapor pressure of the

mixture could not be greater than that of the component with the highest vapor pressure, or less than that of the component with the lowest vapor pressure, and an azeotrope could not occur. Such a system would be ideal, and the non-ideality activity coefficient would be equal to unity. However, only a few binary mixtures follow Kaoult's law at all concentrations. Most systems exhibit either positive or negative deviations from Rapult's law. for those cases where a combination of the magnitude of deviation from kaoult's law, plus the magnitude of the boiling point difference of the two components is such that the vapor pressure of the system will be greater than that of the component with the highest vapor pressure. or less than that of the component with the lowest vapor pressure, an azeotrope will occur. Heterogeneous azeotropes tend to form when the deviations are of such magnitude that immiscible liquid phases are formed.

GREERAL EFFECTS OF PRES UND AND TEMPERATURE ON AZEOTROPES Effect of Pressure and Temperature on Composition

In examining the experimental data of binary azeotropic systems, it can be seen that the composition changes with a change in pressure and temperature. The nature of this change in composition is indicated by Wrewski's rule which can be stated as follows:*

"When the boiling point of a positive azeotrope rises, its composition changes in favor of the component with the higher molecular latent heat of vaporization. --- Conversely, in the case of negative azeotropes, when the boiling point rises, there is an increase in the concentration of the component with the lower latent heat of vaporization."

A graphical representation of this rule is given for a positive azeotrope by Swietoslawski (21). It is pointed out by Swietoslawski, that the upper limit in the increase of pressure (temperature) is the critical point of one of the components. Some azeotropes are known to become non-aseotropic before this point is reached, and others are known to be azeotropic up to the critical point of the components. Experimental data for the ethyl alcohol-water azeotrope shows that this system follows Wrewski's rule (see Table No. 22).

^{*}Perry, J. H., ed., Chemical Engineers' Handbook, Third Edition, p. 631.

The mole fraction of water (the component having the higher molecular latent heat) increases as the pressure (temperature) increases. The data shows that the system becomes non-azeotropic at low pressures. It should be possible theoretically, therefore, to obtain a complete separation of alcohol from water by means of a simple fractionation using a pressure at which the system is non-azeotropic (16).

Pressure-remperature Helationship

in a study of the Vapor pressure-temperature relationship of azectropes, Licht and Denzler (8) found that
equations analogous to the Clapeyron equation should
apply to all types of azectropes. They found that
azectropes have vapor pressure-temperature curves that
are similar to those of pure components, and extend
over the pressure range of the azectrope for a particular system.

Olapeyron equation as a basis, and the fact that the composition of the liquid phase is the same as that of the vapor phase (x1=y1) for an azeotropic system in equilibrium, derived a series of equations that show straight line relationships on a logarithmic plot. In one set of equations the variables are partial pressures, compositions (of either vapor or liquid), and activity coefficients,

versus the total pressure of the azeotrope. In another set of equations the variables are total pressure, partial pressures, compositions (of either vapor or liquid), and activity coefficients, versus the vapor pressure of a pure component. The slopes of the equations involve latent heat ratios and heats of solutions. However, these equations do not exactly define the azeotropic system because the heat ratios were assumed constant in their derivations in order to allow integration of the basic differential equation.

The Cox chart, which is a plot of the Antoine equation, has been used by Mutting and Horsely (12) to predict the azeotropic vapor pressure-temperature curve from two points of azeotropic data, or from the average slope of the pure component vapor pressure curves and one point, of azeotropic data. All three curves are represented as a straight line on the Cox chart.

 $\log P = A - B/(t \neq 230)$ Antoine equation 1 where:

P = pressure.

t = temperature. Ou.

A = constant.

B = constant.

It is shown by Pennington (14), however, that one Antoine line on a Cox chart can only represent one

azeotropic composition at some temperature, but not all azeotropic compositions at once. He therefore objects to the idea that an azeotrope can be represented by a straight line. He indicates a method, in his paper, for determining the true azeotropic curve by the use of a series of Antoine lines, each one representing one azeotropic composition at some temperature. He states that the slope of such a curve can be used to calculate the true molar latent heat of vaporization when substituted in the Clausius-Clapeyron equation.

PUBLISHED METHODS FOR PREDICTING THE EFFECT OF PRESSURE AND TEMPERATURE ON THE COMPOSITION OF AZEOTROPIC SYSTEMS

A number of methods for predicting the effect of pressure and temperature on the composition of azeotropic systems can be found in literature. Some of the methods are based on theoretical considerations, and others have been derived empirically. Several of these methods are reviewed in this section.

The modified Ulausius-Clapeyron equation. mentioned earlier in this paper. Uthmer and Ten Eyck (13) have derived a group of equations that show straight line relationships on logarithmic plots when certain properties of azeotropic systems are plotted against others. used the Clausius-Clapeyron equation as a basis, and also. the condition that $x_1 = y_1$ for an exectropic system in equilibrium. The ratio of the heat quantities was assumed constant in the derivation. One of the equations derived is represented by Equation 2, which gives the relationship of an azeotropic system in terms of composition and azeotropic pressure as a straight line on a logarithmic plot.

$$\log y_1 = (L_1/L_2-1) \log P_2 \neq 0$$

where:

y, = composition of component 1 in vapor phase (mole or wt. %).

Li = molal latent heat of component 1 from solution.

L = molal latent heat of azeotropic solution.

P'z = total pressure of azeotrope.
C' = constant.

= constant.

ascotropic pcints at two pressures, or one ascotropic point and heat data (slope), will completely define an ascotropic system. The disadvantage of this equation is that ascotropic data at more than one pressure, and heat data, are not usually available. Othmer and Ten Eyck point out that the error involved, in the assumption that the heat ratio is constant, would be small, since the change in partial heat of solution of the pure component with concentration change is extremely small compared to the latent heat of vaporisation of the pure component.

A number of asectropic systems are plotted on a logarithmic plot by Othmer and Ten Eyok, and the experimental points are shown to fit very well on a straight line.

Equation based on the two-suffix Margules equation.

Another method for predicting the composition of azeotropic systems with a change in pressure is given by Equation 3.

$$1/x_1 = \left(\frac{\log P_z/P_1}{\log P_z/P_2}\right)^{1/2} \neq 1$$

where:

x₁ = concentration of the more volatile component in the liquid phase (mole fraction).

P. = total pressure of ageotrope.

P₁ = vapor pressure of pure component 1 at azeotrope boiling point.

P₂ = vapor pressure of pure component 2 at asectrope boiling point.

Equation 3 was derived from the symmetrical two-suffix Margules equation for calculating activity coefficients. It may be used for estimating the composition of binary azeotropes when only the total pressure of the azeotrope and the vapor pressures of the pure components at the boiling point of the azeotrope are known. An accuracy of better than $\frac{1}{2}$ 10% is claimed for this method (16).

Method of Carlson and Colburn. A method for predicting the composition of a homogeneous assectrope with change in temperature, that is based on Equation 4 and the property of homogeneous assectropes that $x_1 = y_1$, is given in a paper by Carlson and Colburn (3).

$$P y_1 = r_1 P_1 x_1$$

$$P = r_1 P_1 = r_2 P_2$$
48

$$r_1/r_2 = P_2/P_1$$
 4b

where:

P = total pressure of the system.

P1 = vapor pressure of pure component 1.

Po = vapor pressure of pure component 2.

r; = non-ideality activity coefficient for component 1.

rg = non-ideality activity coefficient for component 2.

x1 = concentration of component 1 in the liquid

phase (mole fraction or mole percent).

y₁ = concentration of component 1 in the vapor phase (mole fraction or mole percent).

The relationship 4b is developed from Equation 4 for the

condition that $\mathbf{x}_1 = \mathbf{y}_1$. If a plot is made of $\mathbf{r}_1/\mathbf{r}_2$ versus the mole fraction of the most volitile component, and another is made of P_2/P_1 versus temperature, both on the same scale, then azeotropes can be picked off where the ordinates for the two curves are equal. This method can be used when activity coefficient and vapor pressure data are available for the pure components of a system.

When activity coefficient data are not available, they can be estimated from azeotropic data at one temperature using either the Margules or the Van Laar equations. (Carlson and Colburn have found that better results are obtained with the Van Laar equations than with the Margules equations (3).) The Van Laar constants are determined from the azeotropic composition data at one temperature, and are then substituted back into the Van Laar equations to estimate the activity coefficients at other concentrations. Since the activity coefficient data are obtained from data at one temperature, some error will be involved in extrapolating to other temperatures. This method will be most accurate when applied to systems where the azeotrope is in the middle half of the composition range.

Azeotropic pressures can then be obtained, from data already calculated, by means of Equation 4s.

The reference point equation. It is suggested by Hougen and Jatson (7) that the azeotropic composition data over a range of temperatures, obtained by the above method of Carlson and Colburn, be used in conjunction with Equation b to calculate azectropic pressures.

$$P_{z} = \frac{P_{z}^{1} (x_{1}P_{1} \neq x_{2}P_{2})}{(x_{1}^{2}P_{1}^{2} \neq x_{2}^{2}P_{2}^{2})}$$

where:

 $P_z = vapor$ pressure of the azeotrope at temperature t. $P_1^z = vapor$ pressure of component 1 at temperature t.

 P_z^1 = vapor pressure of component 2 at temperature t. P_z^1 = vapor pressure of the assotrope at the reference temperature t'.

z vapor pressure of component 1 at the reference temperature t'.

P: = vapor pressure of component 2 at the reference temperature t'.

x1 = mole fraction of component 1 in the exectrope at temperature t.

x2 = mole fraction of component 2 in the azectrope at temperature t.

x; = mole fraction of component 1 in the ageotrope at the reference temperature t'.

xd = mole fraction of component 2 in the azeotrope at the reference temperature t'.

The pressures thus obtained can then be used in Equation 4a to obtain more reliable activity coefficient data. which might be used to obtain a complete equilibrium curve by means of the Margules or Van Laar equations. pointed out that insufficient work along this line has been done to allow an evaluation of the possible errors involved.

Equation b is an empirical equation for predicting azeotropic pressures from azeotropic composition data at different azeotropic boiling points and component vapor pressure data at corresponding temperatures. It is one of the equations tested in this paper against experimental data. The results are given in the next section.

The curves of Horsely. A series of plots for fortyfive systems have been published by Horsely (5) that can be used to estimate boiling point and composition data for azeotropes. These plots were based on equations developed by Lecat for certain related groups of binary systems. Using the plot for the methanol-benzene system. Horsely was able to predict the boiling point of the azeotrope to within f 1 to f 300, and the composition to an accuracy of f 1 to f 4 weight percent, over a range of pressures from 200 to 11,000 Mm. Hg. Boiling point data at different pressures for the pure components are required in order to use the plots. Horsely has found that agreement between predicted and experimental values for most systems is not too good, and suggests that the value of this method lies mainly to serve as a guide in estimating the effect of pressure on asectropes.

The log x equation. A linear relationship for the azeotropic composition as a function of the azeotropic

boiling point has been introduced in a paper by Skolnik (19). The relationship is expressed by Equation 6.

$$\log x = A - BT$$

where:

x = mole percent of one component in the azeotrope.

T = assotropic boiling point (°K).

A = a constant.

B = a constant.

Skolnik shows by a series of plots that this equation represents a number of homogeneous maximum-boiling and hydrocarbon-water heteroazectropes very well. However, the majority of these systems shown have four or less points. In the homogeneous minimum-boiling classification, the methanol-benzene system is shown to be represented fairly well, but the ethanol-water and ethyl acetate-carbontetrachloride azectropes are not represented too well.

the critical points of the components are reached. The point of non-azeotropy would be represented by Equation 6 for a condition when x = 100 mole percent. It would also be the point where the vapor pressure curves of the azeotrope and one of the pure components crossed. Skolnik found this point for the methanol-benzene system solving the Antoine equation for methanol and for the azeotrope.

and checked the corresponding temperature with that obtained from Equation 6 when x = 100 mole percent for methanol. The temperature obtained from the Antoine equations was 202°C, and that obtained from the log x equation was 219°C. Skolnik felt that this was a good check considering the possibility for error in extrapolating the vapor pressure data.

Pennington (14) considered the log x equation as a possibility for representing the l,l-diffuoroethane-dichlorodiffuoromethane system. He rejected it on the grounds that it does not allow x to become zero and therefore cannot truely represent an azeotropic system if Swietoslawski's (21) graphical representation of a maximum-pressure type of azeotrope is correct. Pennington also checked the log x equation against experimental data from four azeotropic systems and found that it did not fit the observed data as well as the T-parabola type of equation which will be considered next.

The T-parabola equation. The T-parabola equation of the type represented by Equation 7 was selected by Pennington (14) to represent the 1,1-diffuoroethane-dichlorodifluoromethane system after rinding that it gave a better fit than several other equations tested against experimental data for this and three other systems.

T = A + Bx + Cx2

7

where:

T = boiling point of the aseotrope (OK).

x = composition of one component in the exectrope (mole %).

A = a constant.

B = a constant.

C = a constant.

The T-parabola equation also allows x to become zero, which allows extrapolation over the entire range of azeotropism for the system using this one equation. However, at least three azeotropic points are required to determine the constants and it is not a linear relationship, therefore, it cannot be applied as readily as the log x equation.

Two of the foregoing equations were selected for testing against experimental azeotropic data, Equations 5 and 6. Equation 5 was selected because no indication of its applicability or adequacy in predicting the pressure-composition relationship for azeotropic systems could be found in the literature. Equation 6 was selected for testing because it appeared to be the most easily applicable, being a linear relationship and requiring only two azeotropic points to define a system. For those systems that become non-azeotropic before the critical points of the components are reached, only one azeotropic

point and the point of non-azeotropy, determined from the antoine equation, are required. Extrapolating over the entire range of the azeotrope is made possible by solving the equation for each component separately, for when one component is 100 mole percent, the other is zero mole percent. In this manner, the points of non-azeotropy can be determined at both ends of the azeotropic system, if such points exist. The work of Skolnik (19) and Pennington (14) indicated that the log x equation represented most azeotropic systems fairly well, and therefore, it was decided to check this equation more extensively.

The results of testing Equations 5 and 6 against published experimental azeotropic data are given in the next section.

TROPIC COMPOSITION WITH CHANGE IN PRESSURE AND TRAPERATURE

The Reference Point Equation (Pressure vs. Composition)

Method of testing. Published data from ten binary azeotropic systems were used in testing Equation b. The systems included seven homogeneous minimum-boiling, two homogeneous maximum-boiling, and one heterogeneous minimum-boiling azeotrope. The azeotropic reference point used in making the calculations was that for atmospheric pressure, or as close to atmospheric as was given in the experimental data. It was felt that this point would be the most accurate since it could be most easily checked against the results of other investigators. Also, it represented a mid-point for most systems. Wherever possible, experimental data from only one investigator were used to avoid the possibility of introducing errors in measurement due to different techniques used by different investigators.

In making the calculations, Equation 5 was simplified to Equation 5a.

$$P_z = K' (x_1 P_1 \neq x_2 P_2)$$
 5a

The vapor pressure data of the pure components were obtained by interpolating and extrapolating from published vapor pressure data using a log P versus $1/({}^{\circ}C \neq 250)$ relationship. The calculations were made with a calculating

machine in conjunction with a five-place logarithm table.

For all systems except the two homogeneous minimumboiling systems, the experimental azeotropic composition
data, and the vapor pressure data for the pure components,
were substituted into the equation and the corresponding
azeotropic pressures were calculated and compared with
the observed values of total pressure. In the case of
the two homogeneous maximum-boiling systems, hydrogen
chloride-water and hydrogen bromide-water, the azeotropic
boiling points were above the critical points of hydrogen
chloride and hydrogen bromide except for the lowest point.
Therefore, no vapor pressure data existed for these two
components in the range of the azeotrope, and the equation
could not be applied. Instead, the equation was revised
to Equation bb and used as such to calculate pseudo vapor
pressure data for these two components.

$$P_1 = (P_2/K' - x_2P_2) / x_1$$
 5b

The reference points used for these systems were the lowest given in the experimental data, since they were the only points below the critical points of the components in question. This pseudo vapor pressure data was then plotted on vapor pressure graph paper along with a plot of vapor pressure data extrapolated from below the critical points of the two components. For the purpose of comparison.

results are presented in Tables 1 and 1a. Tables 2 through 9 give the results for the individual systems tested with the exception of the two homogeneous maximum-boiling systems, to which the equation (Equation 5) did not apply. Figures 1 and 2 represent the results obtained for these two systems using Equation 5b.

In the homogeneous minimum-boiling group, Equation 5 predicted the azeotropic pressures to an overall accuracy of \angle 3% in all systems except the methanol-benzene system. For this system the equation's accuracy was of the order of \angle 15% overall.

There was a trend for the equation to become less accurate as the pressure increased, or decreased, from the reference pressure. For those systems where the overall accuracy was \$\frac{1}{2}\%\$, the deviations were as high as \$\frac{1}{2}\%\$ at pressures of one atmosphere distant from the reference pressure. An exception to this was the n-propanol-water system, for which the percent deviation steadily decreased as far as seven atmospheres away from the reference pressure. The methanol-benzene system also showed the general trend of increasing percent deviation with increase, or decrease, in pressure from the reference pressure. The fact that it showed so great a deviation

is probably due in part to the fact that this system was tested over a much wider span of pressures than any of the other systems. A contributing factor to this trend is the probable reduced accuracy of vapor pressure data at low and high pressures. A straight line relationship was used to extrapolate the vapor pressure data, and the vapor pressure curves are probably non-linear at very low pressures. At high pressures, a slight error in the data used for extrapolation can be magnified a great deal.

The fact that a clear-cut trend for all systems of this type was not obtained probably indicates that errors are present in the experimental azeotropic data. This appears to be especially the case for the methanol-benzene system, for which other experimental data can be found in the literature which do not agree with the data used in this paper (6).

Generally, the results indicate that for homogeneous minimum-boiling azeotropes, Equation 5 will predict the total azeotropic pressure with an accuracy of from \$\frac{7}{2}\%\$ to \$\frac{7}{2}\%\$, within the range of one atmosphere above or below the reference pressure (atmospheric), with the higher deviation occurring as the pressure increases or decreases from the reference pressure. At pressures greater than one atmosphere from the reference pressure (atmospheric), deviations in the magnitude of \$\frac{7}{2}\%\$ to \$\frac{7}{2}\%\$ can be expected.

For the one heterogeneous minimum-boiling system tested, (Table 9, ethyl acetate-water) the results indicate that Equation 5 applies in the same manner as for homogeneous minimum-boiling azeotropes. Although larger deviations were obtained at low pressures (/ 16%) there is no justification for stating that this applies to all systems of this type, since only one such system was tested.

Figures 1 and 2 show that the vapor pressure data for hydrogen chloride and hydrogen bromide, above their respective critical points, obtained by substituting experimental data into Equation 5c, does not coincide with the vapor pressure data for these components extrapolated past their critical points from experimental data below their critical points. Therefore it can be concluded that Equation 5 does not apply to homogeneous maximumboiling systems above the critical points of the components.

The Log x Equation (Composition vs. Temperature)

Method of testing. Published data from ten azeotropic binary systems were used in testing Equation 6. The systems included seven homogeneous minimum-boiling, two homogeneous maximum-boiling, and one heterogeneous minimum-boiling azeotrope. The method of least squares was used for those systems having ten or fewer points, and the method

of averages was used for those systems having more than ten points. The equation was tested for both components of each system.

rindings. The systems used and a summary of the results are presented in Tables 10 and 10a. The results are given for each system separately in Tables 11 through 20. In general it was found that agreement between calculated and observed results was fairly good, less than one mole percent average deviation for a majority of the systems. Equation 6 seemed to represent all types of systems equally well, considering the nature of the experimental data used for each case.

The equation was solved for each component of each system primarily to enable the prediction of non-azeotropic points for the systems. It was found that in most cases better agreement between calculated and observed values was obtained using one component than when the other component was used, when predicting azeotropic compositions. The reason for this is not apparent from the results since no pattern or trend can be detected, and therefore, no conclusions can be drawn as to which component to use in a particular case to obtain the best results (e.g., the most, or least, volatile component, or, the component at the upper, or lower, end of the composition range of the system).

Equations 6 and 7 are compared in Table 21 for their ability to predict non-azeotropic points and to fit experimental data for four systems. The T-parabola type of equation is shown to be superior to the log x type of equation not only when compared to the log x equation as one equation, but also when the other component is represented by x. However, the differences in the results of the two equations are not great. These results agree with Pennington's findings for these two equations (14).

CONCLUSIONS

It has been shown that the reference point equation (Equation 5) can be used to estimate the pressures of binary homogeneous minimum-boiling and heterogeneous minimum-boiling azeotropes, from azeotrope temperature-composition data, with an accuracy of from \$\noting 2\% \to \$\noting 6\% for most systems in the range of one atmosphere pressure above or below the reference pressure (atmospheric). At higher pressures, an accuracy of \$\noting 20\% \to \$\noting 30\% \text{ should be expected. The accuracy of this equation is dependent upon the accuracy of the pure component vapor pressure data used.

It has also been shown that Equation 5 does not apply to binary homogeneous maximum-boiling assotropes in the range above the critical points of the components.

constants for the log x equation (Equation 6) have been calculated and tabulated for both components of ten binary azeotropic systems. It has been shown that the log x equation represents the composition-temperature relationship for these systems very well. Non-azeotropic points of a system can be estimated, when they exist, by solving the log x equation for x = 100 mole percent for each component separately.

A comparrison was made between the log x equation and the T-parabola equation (Equation 7) for their adequacy

in predicting the temperature-composition relationship and non-azeotropic points for four binary azeotropic systems. It was shown that the T-parabola equation gave results that compared more favorably with observed data than did the log x equation, although the differences were not markedly great. However, the T-parabola is not a linear relationship and requires at least three azeotropic points in order to determine the constants. In this respect it is not as easy to apply as the log x equation, which is a linear relationship requiring only two azeotropic points to determine the constants for a system.

In addition, several other methods for predicting the effect of temperature and pressure on the composition of binary assotropic systems. found in the literature, have been reviewed briefly and evaluated on the basis of evidence presented by other investigators.

Finally, the complete experimental data, found in the literature, for the thirteen azeotropic systems used in testing Equations b and 6 have been tabulated and presented in the appendix for easy reference.

TABLE 1. SYSTEMS USED TO TEST EQUATION 6

No.	S.	ystem	Temp. Kange.	Pressure Kange, Mm.	Compo- sition kange, Mole %	Refer ence
	L	(omogeneous	Minimum-B	oiling Azeot	tropes	
1	sthyl water	alcohol-	38 .35- 95 .3 5	94.9- 1451.3	88.69- 100	(11,22)
2	-	acetate- alcohol	-1.37- 91.35	25.0- 1475.5	78.01- 45.13	(10)
3	Methan benzen		26- 149	200-	55.7- 80.7	(5)
4	CC14- ethyl	acetate	47.36- 76.15	285.7- 789.2	55.6- 58.2	(17,18)
5	n-Buta	ne-	-35- 3	549.6- 2014	40.5- 35.6	(9)
6	trans-	2-Butene-	-29- 3	349.6- 1558	27.7- 32.5	(9)
7	n-Prop	-Lons	8 7- 151	740- 5430	43.17- 45.15	(8)
	He	terogeneous	Minimum-	Boiling Azec	trope	
8	Ethyl water	acetate-	-1.90- 89.08	25.0- 1441.3	64.95- 84.56	(10)
	H	omogeneous	Maximum-B	oiling Azeot	ropes	
9	HCl- water		48.724- 122.98	50 - 1220	10.624- 13.126	(2)
10	HBr- water		74.12- 137.34	100- 1200	16.505- 18.091	(1)

^aComposition given is for that of the first component listed.

TABLE 1a. SUMMARY OF RESULTS FOR EQUATION 5

No.	System	Average Deviation Mm. Hg	Average Deviation Percent	No. of Points
	Homogeneous	Minimum-Boiling	z Azeotropes	
1	Ethyl alcohol- water	6.25	3.04	8
2	Ethyl acetate- ethyl alcohol	9. 58	1.70	10
3	Methanol- benzene	1190	15.73	5
4	CCl ₄ - ethyl acetate	6.84	1.87	6
5	n-Butane- SOg	43.40	2.54	5
6	trans-2-Butens- SO ₂	27.88	2.04	4
7	n-Propanol- water	82.5	2.7	5
	Heterogeneo:	us Winimum-Boili	ng Azeotrope	
8	Ethyl acetate- water	15.75	6.27	20
	Homogeneous	Maximum-Boiling	Azeotropes	
9	HCl- water ^a	dingi Milio aper tasp.	wine dips then page	17
10	HBr- water ^a	Note date this vote	aug 400 400	12

aguation 5 does not apply in the range tested.

TABLE 2. PREDICTION OF AZEOTROPE PRESSURE FROM EQUATION 5

IN THE ETHYL ALCOHOL-WATER SYSTEM

No.	Azeotrope Pressure Observeda, Mm. Hg	Asectrope Pressure Calculated, Mm. Hg	Devietion,	Deviation, Percent
1	94.9	100.53	-5.68	5.933
	100.0	104.93	-4.93	4.930
2	129.7	135.73	-6.02	4.641
4	198.4	203.65	-5.25	2.646
5	404.6	412.92	-8.32	2.056
5*	760.0	distribution to the state of the state	-	Marie Santo Agus, maio appo
7	1075.4	1070.01	5.29	0.501
8	1451.3	1443.12	8.18	0.564
AV.	deviation dis	regarding sign	6.25	3. 039

^{*}Reference point.

aReference for asectropic data: (11,22).

TABLE 3. PREDICTION OF AZEOTROPE PRESSURE FROM EQUATION 5

IN THE ETHYL ACETATE-ETHYL ALCOHOL SYSTEM

No.	Azectrope Pressure Observed ^a , Mm. Hg	Azeotrope Pressure Calculated, Mm. Hg	Deviation, Mm. Hg	Deviation Percent
-		es. E. A. HE	0 6 4	0 000
1	25.0	24.43	0.57	2.280
2	77.4	75.95	1.45	1.073
3	117.2	115.16	2.04	1.741
4	219.9	215.15	4.75	2.160
5	423.0	418.19	4.81	1.137
6	578.2	573.57	4.63	0.801
7*	760.0	Note that they died with the	which when made down	
8	948.0	956.27	-8.27	0.872
ÿ	1121.0	1139.07	-18.07	1.612
10	1475.5	1516.79	-41.29	2.798
AV.	deviation disr	egarding sign	9.58	1.697

^{*}Reference point.

akeference for ageotropic data: (10).

TABLE 4. PREDICTION OF AZEOTROPE PRESSURE FROM EQUATION 5

IN THE METHANOL-BENZENE SYSTEM

No.	Azeotro; Pressure Observed Mm. He	Pressure 1 ^a , Calculated,	Deviation, Mm. Hg	Deviation, Percent
1	200	183	17	8.50
2	400	3 9 3	7	1.75
2 3*	76 0	Window Steam opings.		water white shade
4	6,000	7.271	-1271	21.18
5	11,000	14,465	-3465	31.50
Av.	deviation	disregarding sign	1190	15.73

^{*}Reference point.

a Reference for asectropic data: (5).

TABLE 5. PREDICTION OF AZEOTROPE PRESSURE FROM EQUATION 5

IN THE CARBON TETRACHLORIDE-ETHYL ACETATE DYSTEM

No.	Azeotrope Pressure Observed ^a . Mm. Hg	Azectrope Pressure Calculated, Mm. Hg	Deviation, Mm. Hg	Deviation, Percent
1	285.7	288.67	-2.97	1.04
2	385.2	381.85	3.35	0.87
3	484.5	474.18	10.32	2.13
4	583.7	573.82	9.88	1.69
5	685.0	677.30	7.70	1.12
6*	789.2	which which wider their short-	that where now retires	1900-1600 1804-1804
ÀV.	deviation di	regarding sign	6.84	1.37

^{*}Reference point.

a Reference for azeotropic data: (17,18).

TABLE 6. PREDICTION OF AZEOTROPE PRESSURE FROM EQUATION 5

IN THE n-BUTANE-SULFUR DIOXIDE SYSTEM

10.	Azeotrope Pressure Observed Mm. Hg	Pressure B, Calculated,	Deviation, Mm. Hg	Deviation, Percent
1	349.6	346.03	3. 57	1.02
2*	752.4	whose stages women and a digital public	2006 HORE HARE MORE	
3	1436.5	1413.34	23.16	1.61
4	1436.5	1416.02	20.48	1.43
b	2014	2140.39	-126.39	6.28
Av.	deviation (dieregarding sign	43.40	2.59

^{*}Reference point.

akeference for ageotropic data: (9).

TABLE 7. PREDICTION OF AZEOTROPE PRESSURE FROM EQUATION 5

IN THE TRANS-2-BUTENE-SULPUR DIOXIDE SYSTEM

No.	Azeotrop Pressure Observed Ma. Hg	Pressure a, Calculated,	Deviation,	Deviation, Percent
1	349.6	351.93	-2.33	0.67
2*	752.4	-quipe state spice spice appro-		-
3	1428.8	1471.78	-42.43	3.00
1 2* 3 4	1558	1596.37	-36.37	2.46
AV.	deviation	disregarding sign	27.88	2.04

^{*}Reference point.

akeference for asectropic data: (9).

TABLE 8. PREDICTION OF AZEOTROPE PRESSURE FROM EQUATION 5

IN THE n-PROPANOL-WATER SYSTEM

No.	Azeotrope Pressure Observed ^a , Mm. Hg	Azeotrope Preseure Calculated. Mm. Hg	Deviation. Mm. Hg	Deviation, Percent
1* 2 3	740	the with the law.	white states	alon mink signs.
2	1790	1727	63	3.5
3	2830	2 725	107	3.8
4	3860	3770	y 0	2.3
5	5930	5860	70	1.2
AV.	deviation dis	regarding sign-	- 82.5	2.7

^{*}Reference point.

aReference for azeotropic data: (8).

TABLE 9. PREDICTION OF AZEOTROPE TRESSURE FROM EQUATION 5

IN THE ETHYL AGETATE-WATER SYSTEM

	Azeotrope Pressure	Azeotrope Pressure		
No.	Observed ^a .	Calculated.	Deviation. Mm. Hg	Deviation, Percent
1	25.0	29.18	-4.18	16.72
2	50.0	58.10	-8.10	16.20
3	78.5	90.76	-12.26	15.62
4	82.2	95.05	-12.86	15.63
5	150.0	166.09	-16.09	10.73
6	176.0	194.26	-18.26	10.38
7	250.0	268.78	-18.78	7.51
8	329.8	348.13	-18.33	5.56
9	420.0	436.72	-16.72	3.98
10	446.2	442.74	3.46	0.78
11	606.0	613.17	-7.17	1.18
12	613.8	635.99	-22.19	3. 62
13	745.0	746.89	-1.89	0.25
14*	760.0	the six six the six six	the run aire tim	White make water where
15	875.0	871.79	3.21	0.37
16	903.5	896.37	7.13	0.79
17	984.3	971.96	12.34	1.25
18	1177.9	1152.80 -	25.10	2.13
19	1415.0	1369.95	45.05	3.18
20	1441.8	1395.15	46.15	3.20
AV.	deviation disre	egarding sign	15.75	6.27

^{*}Reference point.

aReference for azeotropic data: (10).

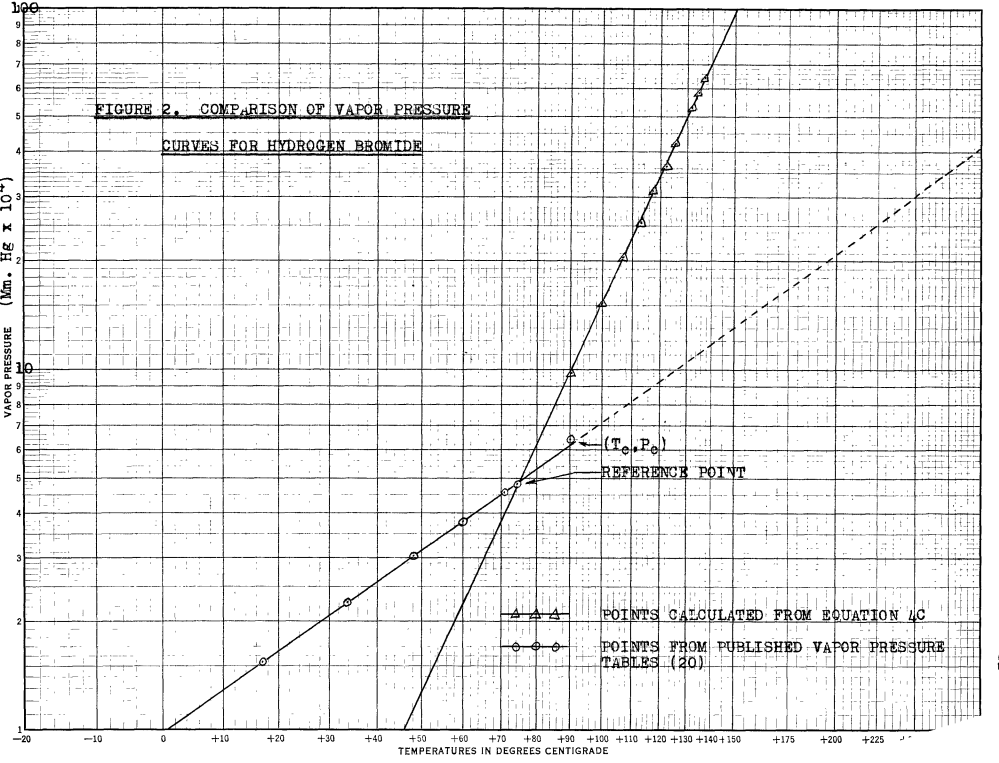


TABLE 10. SYSTEMS USED TO TEST EQUATION 6

No.	System	Temp. Kange,	Fressure Kange, Mm.	Compo- sition Kange ⁸ , Mole %	kefer-
	Homogeneous	Minimum-Bo	iling Azeot	ropes	
1	Ethyl alcohol-	33.36-	94.9-	88.69-	(11,28)
	water	96.35	1451.3	100	
2	Mthyl acetate-	-1.37-		78.01-	(10)
	ethyl alcohol	91. 35	1475.5	45.13	
3	Methanol-	26-	-003	55.7-	(5,19)
	benzene	149	11,000	80.7	•
4	0014-	47.36-		55.6-	(17,18)
	ethyl acetate	76.16	789.2	58.2	
5	n-Butane-	-36-	349.6-	40.5-	(9)
	sog	8	2014	35.6	(
6	Isobutene-	-35-	349.6-	46.0-	(9)
	sog	3	1824	37.5	
7	DCDFMp-	-30.5-	tivity team date, date	65.36-	(14)
	DEE	40.08		55.22	
	Heterogeneous	e Winimum-B	oiling Aseot	rope	
8	Ethyl acetate-	-1.9-	25.0-	84.56-	(10)
	water	89.08	1441.3	64.95	
	<u>Homogeneous</u>	Maximum-Bo	iling Ameot	ropes	
9	HCl- water	48.724- 122.98	50- 1220	13.126- 10.624	(2)
10	HBr-	74.12-	100-	18.091-	(1)
	Water	137.34	1200	16.605	

Composition given is for that of the first component listed.

bucdFM = dichlorodifluoromethane.

Coff = 1,1-difluoroethane.

TABLE 10a. SUMMARY OF RESULTS FOR EQUATION 6

		Oon	stants	Av. Devi- ation.	no. of
No.	System	A	В	Mole %	Points
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Homo geneou	s Minimum-E	oiling Azeot	ropes	
1	sthyl alcohol-	2.22630	0.0007713	0.85	8
	water	-4. 05609	-0.0143062	2.10	
2	Ethyl acetate-	2.48849	0.0021885	1.76	10
•••	ethyl alcohol	0.15963	-0.0043540	0.37	
5	Methanol-	1.3508	-0.0018152	0.56	10
•	benzene	2.62433	0.0028802	1.40	
4	44.57 m. 1	3.07827	0.0380721	0.53	6
*	col ₄ -	-0.87244	-0.0072389	0.74	Ü
_	•	15	and an article and the second and the second		¥
5	n-Butane-	1.99659	0.0016007	0.42	4
	30 ₂	1.55771	-0.0008994	0.4	
6	Isobutene-	2.14299	0.0019337	3.14	6
	30 ₂	1.37539	-0.0014295	3.1	
7	DCDFMp-	2.06353	0.0010246	0.12	4
•	DIE	1.16256	+0.0015656	0.24	-
	Heterogene	ous Minimum	-Boiling Aze	otrope	
8	Ethyl acetate-		0.0013730	0.265	20
O	water	0.12664	-0.0029570	0.68	Shir Sar
	Homopanaon	H-mumirak a	oiling Azeot	robes	
9	HC1-	1.57192	0.0018772	0.046	17
•	water	1.87979	-0.0001807	0.037	yan. F
***	72 * h.c.	n entere	A AAA####	ለ ለደላ	* 4
10	HBr-	1.52875	0.0007596	0.062	12

The constants for both components of each system are given (log x = A - BT, where: x = mole %, T = OK).

b DUDFM = dichlorodifluoromethane.

ODFE = 1,1-difluoroethane.

TABLE 11. DEVIATIONS FROM OBSERVED VALUES IN PREDICTING
AMBOTROPIC COMPOSITIONS FROM EQUATION 6 IN THE
ETHYL ALCOHOL-WATER SYSTEM

	Azeotrope Boiling Point ^a ,	Azeotrope Composition ⁸ , Mole %	Deviationb, Mole %	
No.	°K.	EthoH	x=EthOH	x= H ₂ O
1	306.51	98.73	1.00	-0.86
2	307.36	98.63	1.07	-0.82
3	312.36	y6.74	0.05	0.67
4	320.79	¥3.37	-1.89	3.22
5	336.20	91.94	-0.74	3.39
6	351.31	89.47	-0.76	1.20
7	360.28	88.91	0.10	-1.45
8	368.51	88.69	1.18	-5.20
AV.	deviation di	eregarding sign	0.85	2.10

aReference for azeotropic data: (11,22).

Deviations have a positive sign when the calculated value is smaller.

TABLE 12. DEVIATIONS FROM OBSERVED VALUES IN PREDICTING

AZEOTROPIC COMPOSITIONS FROM EQUATION 6 IN THE

ETHYL ACETATE-ETHYL ALCOHOL SYSTEM

	Azeotrope Boiling Point ^a .	Azeotrope Composition ^a , Mole %	Deviation ^b , Mole %	
Vo.	o _K .	Ethac	x= sthac	x= sthOH
1	271.79	78.01	-0.27	-0.04
2	291.87	73.37	2.62	-0.31
3	300.18	71.00	2.15	-0.89
4	313.66	86.02	2.63	0.46
5	324.47	60.0H	1.63	0.64
6	337.59	56.85	0.66	0.54
7	344.97	53.87	-0.27	0.25
8	351.29	51.12	-1.38	0.00
ÿ	356.21	49.03	-2.13	-0.38
LO	364.51	45.13	-3.98	-0.81
. v. e	leviation dis	regarding sign	1.76	0.37

a Reference for azeotropic data: (10).

Deviations have a positive sign when the calculated value is smaller.

TABLE 18. DEVIATIONS FROM OBSERVED VALUES IN PREDICTING
AZEOTROPIC COMPOSITIONS FROM EQUATION 6 IN THE
METHANOL-BENZENE SYSTEM

	Azeotrope Boiling Point	Azectrope Composition ^a , Mole %	Deviation ^b , Mole %	
No.	o _K .	Methanol	x=Methanol	x=Benzene
1	299.16	55.7	0.20	-2.23
2	298.16	55.6	0.27	-2.40
3	315.16	57.5	-0.75	0.66
4	313.16	58.7	0.80	-1.10
5	329.86	59.9	-1.00	2.14
6	329.86	59.9	-1.00	2.14
7	371.46	61.5	0.30	0.95
8	330.16	61.9	0.94	0.22
y	397.16	74.8	0.13	0.91
10	422.16	80.7	0.16	-1.28
AV. C	eviation dis	regerding sign	0.56	1.40

aReference for assotropic data: (5,19).

b Deviations have a positive sign when the calculated value is smaller.

AZEOTROPIC COMPOSITIONS FROM EQUATION 6 IN THE

CARBON TETRACHLORIDE-ETHYL ACETATE SYSTEM

	Azeotrope Boiling Point	Azeotrope Composition ^a , Mole %	Deviation ^b . Mole %	
No.	<u>°ж.</u>	0014	x=0014	x=EthAc
1	320.52	72.60	0.48	-1.64
2	328.38	67.75	1.98	0.29
3	334.48	63.73	-0.07	0.89
4	339.88	60.75	-0.10	0.53
5	344.72	58.2	-0.12	-0.19
6	349.31	55.60	-0.42	-0.91
AV.	deviation aid	regarding sign	0.58	0.74

aReference for asectropic data: (17,18).

bDeviations have a positive sign when the calculated value is smaller.

TABLE 15. DEVIATIONS FROM OBSERVED VALUES IN PREDICTING

AZEOTROPIC COMPOSITIONS FROM EQUATION 6 IN THE

n-BUTANE-SULFUR DIOXIDE SYSTEM

	Azeotrope Boiling Pointe,	Azeotrope Composition ^a , Nole %	Deviation ^b , Mole %	
ro.	o _K .	n-Butane	x=n-Butane	x=80g
1	238	40.5	-0.77	0.4
2	255	39 . 3	0.54	-0.5
3	255	28.9	0.14	-0.1
4	254	39.7	0.79	-0.8
5	268	36.9	-0.05	0.2
6	268	36.6	-0.35	0.5
7	276	35.6	-0.28	0.4
AV.	deviation di	eregarding sign	0.42	0.4

aReference for assotropic data: (9).

b Deviations have a positive sign when the calculated value is smaller.

AZEOTROPIC COMPOSITIONS PROM EQUATION 6 IN THE

ISOBUTENE-SULFUR DIOXIDE SYSTEM

	Azeotrope Boiling Foint ^a ,	Aseotrope Composition ^a , Nole %	Deviation ^b , Mole %	
No.	ox.	Isobutene	x=lsobutene	x=SO ₂
1	243	46.0	-1.11	1.2
2	259	44.2	0.33	0.0
3	273	45.6	4.38	-3.9
4	273	46.4	5.18	-4.7
5	276	36	-4.67	5.0
6	276	37.5	-3.17	3.5
AV.	deviation dis	eregarding sign-	- 3.14	3.1

Reference for assotropic data: (9).

b Deviations have a positive sign when the calculated value is smaller.

TABLE 17. DEVIATIONS FROM OBSERVED VALUES IN PREDICTING
AZEOTROPIC COMPOSITIONS FROM EQUATION 6 IN THE
DICHLORODIFLUOROMETHANE-1,1-DIFLUOROETHANE SYSTEM

	Aseotrope Boiling Point ^a .	Azeotrope Composition ^a , Mole %	Deviation ^b , Nole %	
No.	o _K .	DODPid	x=DCDFM	x=DPB
1	242.66	65.36	0.06	-0.23
2	273.16	60.60	-0.17	0.47
3	298.06	57.47	0.17	-0.05
4	313.24	55.22	-0.06	-0.20
AV.	deviation di	eregarding sign	0.12	0.24

BReference for azeotropic data: (14).

bDeviations have a positive sign when the calculated value is smaller.

TABLE 18. DEVIATIONS FROM OBSERVED VALUES IN PREDICTING
AZEOTROPIC COMPOSITIONS FROM EQUATION 6 IN THE
ETHYL ACETATE-WATER SYSTEM

	Azeotrope Boiling Point ^a ,	Azeotrope Compositiona, Devia Mole % Mole		
No.	o _K .	EthAe	x=Sthac	x≠H20
1	271.26	84.56	-2.06	-0.41
2	283.21	83. 22	-0.19	-0.89
3	291.61	81.49	0.76	-1.07
4	292.54	81.63	0.64	-0.87
5	304.51	78.21	0.23	0.33
6	307.98	77.51	0.38	0.34
7	315.71	75.32	0.05	0.92
8	322.22	73.55	-0.19	1.23
9	328.10	72.38	0.00	1.02
O	329.60	72.26	0.22	0.77
.1	337.49	70.42	0.16	0.60
.2	337.76	70,22	0.02	0.73
13	342.99	69.18	0.13	0.35
.4	343.54	68.85	-0.08	0.52
.5	347.54	68.05	-0.01	0.19
.6	348.39	67.78	-0.10	0.21
17	350.82	67.38	0.02	-0.10
18	356.11	66.18	-0.06	-0.52
.9	361.65	65.05	-0.04	-1.17
0:	362.24	64.95	-0.02	-1.26
iv. d	eviation dis	regarding sign	0.265	0.68

aReference for azeotropic data: (10).

bDeviations have a positive sign when the calculated value is smaller.

TABLE 19. DEVIATIONS FROM OBSERVED VALUES IN PREDICTING
AZEOTROPIC COMPOSITIONS FROM EQUATION 6 IN THE
HYDROGEN CHLORIDE-WATER SYSTEM

	Azeotrope Boiling Point ^a ,	Azeotrope Compositiona, Mole %	De viation^b, Mole %	
io.	°K.	кој	x=H0l	x= H ₂ 0
1	321.884	13.126	-0.521	0.189
1 2	343.116	12.557°	-0.014	-0.010
3	354.365	12.158	0.027	-0.022
4	363.397	11.870°	0.082	-0.064
5	365.240	11.754 ⁸	0.034	-0.017
6	368.189	11.655°	0.045	-0.026
7	370.738	11.557	0.040	-0.020
8	372.813	11.4710	0.030	-0.011
9	375.369	11.385°	0.036	-0.020
.0	377.127	11.305	0.019	-0.005
11	378.724	11.247	0.018	-0.005
.2	379.584	11.214°	0.016	-0.005
13	381.019	11.158	0.010	-0.001
4	381.744	11.130	0.008	0.001
15	383.167	11.089	0.017	-0.011
16	289.545	10.831	-0.026	0.017
.7	395.14°	10.624	-0.035	0.009
. v. đ	leviation dis	egarding sign	0.046	0.057

⁸Reference for azeotropic data: (2).

bDeviations have a positive sign when the calculated value is smaller.

c Interpolated.

TABLE 20. DEVIATIONS FROM OBSERVED VALUES IN PREDICTING

AZEOTROPIC COMPOSITIONS FROM EQUATION 6 IN THE

HYDROGEN BROWIDE-WATER SYSTEM

	Azeotrope Boiling Point ^a ,	Azeotrope Composition ^a , Nole %	Deviation ^b , Mole %	
No.	o _K .	HBr	x=HBr	x•iig0
1	347.28	18.091	-0.302	1.182
2	363.51	17.786	-0.106	0.385
3	373.07	17.523	-0.071	-0.009
4	380.16	17.216	-0.061	-0.293
5	386.10	17.156	-0.012	-0.546
6	390.98	17.020	-0.032	-0.751
7	395.16	16.901	-0.104	-0.926
8	898.95	16.800	-0.019	-190
មួ	402.29	16.710	-0.007	-1.236
10	405.28	16.638	0.008	-1.375
11	407.96	16.566	0.012	-1.494
12	410.50	16.505	-0.012	-1.614
LV. đ	eviation dis	regarding sign	0.062	0.909

aReference for azeotropic data: (1).

bDeviations have a positive sign when the calculated value is smaller.

TABLE 21. A COMPARISON OF THE LOG X AND THE T-PARABOLA

EQUATIONS FOR ADEQUACY IN PREDICTING THE TEMPERATURE—

COMPOSITION RELATIONSHIP AND NON-AZEOTROPIC POINTS OF

AZEOTROPIC SYSTEMS

	4.4	Com-	Av. Devi-	remp. at azeotrop	y. OK.	
System	Method Used	ponent (x)	ation,		At High Pressure	Refer-
070004	VOOL		MOIO 10	TIOSECTA	tragamia	ance.
DCDFMb-	log x	DFE	0.24	abote which regard enter relate	534.9	
DFEO	log x	DCDFM	0.12	63	-	
	T-parabola	DFE	0.11	112	967	(14)
EthOH d-	log x	EthAc	1.76	223.2	shift when more takes where	
EthAce	log x	EthOH	0.44	-	423.4	
	T-parabola	BthoH	0.24	184.4	401.1	(14)
BthOH-	***	EthOH	0.85	apply when rathe space	423.3	
water	log x	water	2.10	243.4		
	T-parabola	water	April 2005 APRIL 2009	314.0	665.3	
	Experiments	a 1	bijer dada sirih silika	307	Marke affiliar scane waves Marke	(14)
MeoH ^f -	log x	Benzene		183.8	then pair facts then apply	
Benzene	log x	MeOH	0.56		493.6	14,19)
	T-parabola	Meoh	0.56	Neg. value	481	(14)
	Antoine	HOOM		Mark made 1974 hands appear	475	(19)

a where no reference is given, the data was calculated by the Author.

b DCDFM = dichlorodifluoromethane.

OFFE = 1,1-difluoroethane.

dsthOH = ethyl alcohol.

e EthAc = ethyl acetate.

f MeOH = methyl alcohol.

APPENDIX

TABLES

of

PRESSURE-TEMPERATURE-COMPOSITION DATA

FOR

BINARY AZEOTROPES

AND

PHYSICAL PROPERTIES OF THEIR COMPONENTS

TABLE 22

	4/	the state of the s		omponent 1	Component 2
Syst	em		-	nyl Alcohol	Water
Form	ula			C ₂ H _b O	H2O
Form	ula We	ight		46.07	18.016
Norm	al Boi	ling Poin	t, ^o c.	78.4	100.0
Melt	ing Po	int, oc.		-112	0.0
Orit	ical Te	mperatur	e, ^o ⊄.	243.5	374.2
Crit	ical Pr	essure.	m. Hg	47,956	165,6 80
	-			refigieure - commente de la president de la constanta de la constanta de la constanta de la constanta de la co	
	oiling	zeotropio	o Data ^a Composition,	· Vones Dec	
J.	oint.	Pressure			esure of Pure
No.	orne,		Mole % Component 1	Compone	nts, Am. Hg
No.		Mm. Hg 70.0 94.9	Mole % Component 1 100.00 98.73	Component Component 94.7	onts, Am. Hg 18 Component 28
No. 1 2 3 4	33.35 34.2 39.20	70.0 94.9 100.0 129.7	Mole % Component 1 100.00 98.73 98.63 96.74	Compone Component 94.7 98.9 129.4	28.0 39.9 52.7
No. 1 2 3 4 5	33.35 34.2 39.20 47.63	70.0 94.9 100.0 129.7 198.4	Mole % Component 1 100.00 98.73 98.63 96.74 93.37	Component 94.7 98.9 129.4 198.1	28.0 39.9 52.7 81.7
No. 1 2 3 4 5	33.35 34.2 39.20 47.63 63.04	70.0 94.9 100.0 129.7 198.4 404.6	Mole % Component 1 100.00 98.73 98.63 96.74 93.37 91.94	Component 94.7 98.9 129.4 198.1 403.0	28.0 38.0 39.9 52.7 81.7 171.2
No. 1 2 3 4 5 6	33.35 34.2 39.20 47.63 63.04 78.15	70.0 94.9 100.0 129.7 198.4 404.6 760.0	Mole % Component 1 100.00 98.73 98.63 96.74 93.37 91.94 89.47	Compone Component 94.7 98.9 129.4 198.1 403.0 755.4	28.0 28.0 39.9 52.7 81.7 171.2 329.0
No. 1 2 3 4 5 6 7 8	33.35 34.2 39.20 47.63 63.04	70.0 94.9 100.0 129.7 198.4 404.6	Mole % Component 1 100.00 98.73 98.63 96.74 93.37 91.94	Component 94.7 98.9 129.4 198.1 403.0	38.0 39.9 52.7 81.7 171.2

aReference: (11,22).

TABLE 23

		and the later of the second and the second control of the second
	Component 1	Component 2
System	Ethyl Acetate	Ethyl Alcohol
Formula	C4H8O2	с ₂ н ₆ 0
Formula Weight	88.10	46.07
Normal Boiling Point, OC.	77.1	78.4
Melting Point, OC.	82.4	-112
Critical Temperature, °C.	250.1	243.5
Critical Pressure, Mm. Hg	28,804	47,956

V/(-m-part)		Azeotropio	Data ^a		
	Boiling Point,	Preseure	Composition,	Vapor Press Component	ure of Pure s. Mm. Hg
No.	. °C.	Mn. Hg	Component 1	Component 18	Component 2
1	-1.37	25.0	78.01	22.2	10.7
2	18.71	77.4	73.37	68.7	40.4
3	27.02	117.2	71.00	105.5	56.4
4	40.50	219.9	66.02	191.3	138.2
5	56.31	423.0	60.08	362.4	298.2
6	64.43	578.2	56.85	487.7	427.9
7	71.81	760.0	53.87	635.0	585.2
8	78.13	948.0	51.12	784.7	754.8
9	83.05	1121.0	49.05	920.4	914.3
10	91.35	1475.5	45.13	1190.8	1246.7

aReference: (10).

bReference: (11).

TABLE 24

				Component 1	Component 2
Sys	System			Methanol	Benzene
For	mula			CH ₄ O	с ₆ н ₆
For	mula We:	lght		32.04	78.11
Nor	mal Boi	ling Point,	°C.	64.7	80.1
Mel	ting Po	int, oc.		-97.8	5.5
Cri	tical To	emperature,	°C•	240.0	290.5
77-4	tinol Di	ressure. Mo	59,812	38,076	
VII	n-recore *			*	•
V.F.I.	brock 13				
V.F.I.		Aseotropic		nagadangiya dan addinadiga addin angalaniyya dan edib oo kalaadii addin dan dan a	
***************************************	Boiling	Assotropic C	Data ^a Compositio	on, Vapor Pre	seure of Pure
***************************************	Boiling Point,	Azeotropio C Pressure,	Data ^a Composition	on, Vapor Pre	nte. Mm. Hg
***************************************	Boiling Point,	Assotropic C	Data ^a Composition	on, Vapor Pre	
No.	Boiling Point,	Asectropic C Pressure, Mm. Hg C	Data ^a Composition Mole % Component	on, Vapor Pre Compone Component	nts, Mm. Hg 1 ^b Component 2 ^b
No.	Boiling Point, °C.	Assotropic Pressure, Mm. Hg 0	Data ^a Composition Mole % Component	On, Vapor Pre Compone Component 1 Component	nts, Mm. Hg 1 Component 2 Properties 195.03
No.	Boiling Point, °C. 26 25	Azeotropic Pressure, Mm. Hg C	Data ^a Composition Mole % Component 55.7 55.6	On, Vapor Pre Compone Component 126.9 120.4	nte, Mm. Hg 1 ^b Component 2 ^t 95.03 90.7
No.	Boiling Point, OC. 26 25 42	Assotropic Pressure, Mm. Hg 0	Data Composition Mole % Component 55.7 55.6 57.5	Vapor Pre Compone Component 126.9 120.4 279.0	nts. Mm. Hg 1 ^b Component 2 ^b 95.03 90.7 191.9
No. 1 2 3 4	Boiling Point, °C. 26 25 42 40	Pressure, Mm. Hg 0 200 223 400	Data ^a Composition Mole % Component 55.7 55.6 57.5 58.7	Component 1 Component 126.9 120.4 279.0 254.2	nts. Mm. Hg 1
No. 1 2 3 4 5	Boiling Point, °C. 26 25 42 40 56.7	Pressure. Mm. Hg 0 200 223 400 725.5	Data ^a Composition Mole % Component 55.7 55.6 57.5 58.7 59.9	1 Component 1 Component 126.9 120.4 279.0 254.2 537.2	nts, Mm. Hg 1
No. 1 2 3 4 5 6	Boiling Point, °C. 26 25 42 40 56.7 56.7	Azeotropic Pressure, Mm. Hg 0 200 223 400 725.5 737	Data ^a Composition Mole % Component 55.7 55.6 57.5 58.7 59.9	Component 1 Component 126.9 120.4 279.0 254.2	nts. Mm. Hg 1
No. 1 2 3 4 5 6 7	Boiling Point, °C. 26 25 42 40 56.7 56.7 58.3	200 223 400 725.5 737	Data Composition Mole % Component 55.7 55.6 57.5 68.7 59.9 61.5	200, Vapor Pre Compone 1 Component 126.9 120.4 279.0 254.2 537.2 537.2	nts. Mm. Hg 1
No. 1 2 3 4 5 6	Boiling Point, °C. 26 25 42 40 56.7 56.7	Azeotropic Pressure, Mm. Hg 0 200 223 400 725.5 737	Data ^a Composition Mole % Component 55.7 55.6 57.5 58.7 59.9	1 Component 1 Component 126.9 120.4 279.0 254.2 537.2	nts, Mm. Hg 1

aReferences: (5,19).

bReference: (5).

TABLE 25

			<u>C</u> c	omponent 1	Component 2
Sys	tem		Carbon	Tetrachloride	Ethyl Acetate
For	mula			CO1 ₄	° ₄ н ₈ ° ₂
For	mula We	ight		153.84	88.10
Nor	mal Boi	ling Point	·,°0.	76.75	77.1
Mel	ting Po	int, °C.		-22.6	-82.4
Ori	tical T	emperatur	9,°¢.	283.1	250.1
Cri	tical P	resaure, l	im. Iig	34,200	28,804
		and the state of t			the second section of the sect
		Aseotropic		4-1	-
	Boiling Point.	Pressure	Composition Mole %	·	sure of Pure
No.	٥٥.	Mm. Hg	Component	Component 1	p Component Sp
1	47.36	285.7	72.60	279.36	250.21
2	55.22	385.2	67.75	366.93	342.26
3	61.32	484.5	63.73	453.66	431.93
	66.72	583.7	60.75	547.06	527.66
4	WU # I G	44 m 35 m 1			
	71.56	685.0	58.2	643.57	627.46

aReferences: (17,18).

bReference: (20).

TABLE 26

	Component 1	Component 2
System	n-Butane	Sulfur Dioxide
Formula	c ₄ H ₁₀	soz
Formula Weight	58.12	64.06
Normal Boiling Point, o	C0.6	-10.0
Melting Point, OC.	-136	-73.2
Critical Temperature, o	C. 152.8	157.2
Critical Pressure, Ma.	Hg 27,360	59,052
Asectropie Da	ta ^a	
Boiling Com	position, Vapor	Pressure of Pure
No. OC. Mm. Hg Com	ponent 1 Uompone	nt 1 Component 2
1 -35 349.6	40.5 164.8	9 204.80
2 -18 752.4	39.3 294.7	5 516.8 0
3 -18 752.4	38.9 294.7	3 516.80
4 -19 752.4	39.7 286.6	8 491.46
5 -5 1436.5	36.9 463.2	2 950.14
	36.6 463.2	2 950.14
	35.6 869.1	2 1831.37

Reference: (9).

bReference: (20).

TABLE 27

	Component 1	Component 2
System	trans-2-Buten	e Sulfur Dioxide
Formula	c₄H8	30 ₂
Formula Weight	56.10	64.06
Normal Boiling Point, OC.	1.0	-10.0
Melting Point, °C.	-105.4	-73. B
Critical Temperature, OC.	sinja man-sinan 4983	157.2
Critical Pressure, Mm. Hg	MAC date date 400	59,052
Azeotropic Data		
		Preseure of Pure
Point, Pressure, Mole	nent 1 Compon	ponents, Mm. Hg ent 18 Component 2
1 -29 349.6 32.	5 207.	10 289.09
2 -14 752.4 81.	•	

27.8

27.7

762.88

822.72

1226.37

1331.37

1

4 3

1428.8

1558

akeference: (Y).

bReference: (20).

TABLE 28

		Com	ponent 1	Component 2	
System		Is	obutene	Sulfur Dioxid	.e
Formula			C₄H႘	sog	
Pormula F	Veight		56.10	64.06	
Normal Bo	oiling Point	t. oc.	-6.7	-10.0	
Melting I	Point, oc.		Willy sale fine and	-73.2	
Oritical	remperature	o. oc.	STOP TRANS MISS AND .	157.2	
Oritical	Pressure,	un. Hg	quide distant stillar sales	59,052	
	Azeotropio) Data ⁸			-
Boiling Point,	Azeotropio	Data ^a Compositi	P 95	by,0b2 Pressure of Pur ponents, Am. Hg	· e
Boiling Point,	Azeotropio E Pressure	Data ^a Compositi	<u> Čom</u>	Pressure of Pur	-
Boilin Point,	Azeotropio E Pressure	Data ^a Compositi Mole %	<u> Čom</u>	Pressure of Pur	2
Boiling Point,	Azeotropio E Pressure, Mm. Hg	Data ^a Compositi Mole % Component	<u> Čom</u>	Pressure of Pur ponents, Am. Hg snt 1 Component	2
Boiling Point, No. Oc.	Azestropio B Pressure Mm. Hg 349.6	Data ^a Compositi Mole % Component	<u> Čom</u>	Pressure of Pur ponents, Am. Hg ant 1 Component	
Boiling Point, No. °C.	Azeotropio Pressure Am. Hg 349.6 752.4	Data ^a Compositi Mole % Component 46.0 44.2	<u> Čom</u>	Pressure of Pur ponents, Am. Hg ent 1 Component 273.34 628.94	2
Boiling Point, No. °C.	Azeotropio Bressure Lim. Hg 349.6 752.4 1428.8	Data a Compositi Mole % Component 46.0 44.2 45.6	<u> Čom</u>	Pressure of Pur ponents, Am. Hg ent 1 Component 273.34 628.94	2

aReference: (9).

b_{Reference: (20).}

TABLE 29

	Component 1	Component 2
System	n-Propanol	Water
Formula	o ₃ H ₈ o	H ₂ O
Pormula Weight	60.09	18.016
Normal Boiling Point, Ou.	97.8	100.0
Melting Point, oc.	-127	0.0
Critical Temperature, oc.	263.7	374.2
Critical Pressure, Mm. Hg	37,924	165,680

		sectropic) Data ^a		
	Boiling Point.	Pressure	Composition, Mole %	***	oure of Pure
No.	°0.	Mm. Hg	Component 1	Component 1	Component 2
1	87	740	43.17	493.5	468.7
2	110	1790	43.78	1191.3	1074.56
3	124	2830	44.07	1885.6	1687.81
4	135	3860	44.52	2545.8	2347.26
5	151	5980	45.15	4006.1	3667.00

aReference: (8).

bkeference: (16).

Ckeference: (4,15).

TABLE 30

	Component 1	Component 2
System	DODFM	υFE ^b
rormula	rgcclg	c ₂ H ₄ F ₂
Formula Weight	120.92	66.05
Normal Boiling Point, OC.	-29.2	-24.7
Melting Point, °C.	-156	
Critical Temperature, °C.	111.6	108
Critical Pressure, Mm. Hg	30,096	style title allen dele

		Aseotropic) Data ⁰		
	Boiling Point,	Pressure,	Composition,	Component	ure of Pure
No.	. 00.	Mm. Hg	Component 1	Component 1	Component 2
1	-30.50	**** **** ****	65.36	then aims rein 400	allette fander statet
2	0.00		60.60	-	-
3	24.90	-	57.47	-	-
4	40.08	the state only with	55.22	Title sain title dide	Apper value value guals.

aDCDFM = dichlorodifluoromethane.

bDFE = 1,1-difluoroethane.

CReference: (14).

TABLE 31

		Compor	ent 1	Component 2
System	•	Ethyl A	oetate	Water
Formula		0 ₄ H ₈	02	H ₂ O
Formula Weig	ght	88.	10	18.016
Normal Boili	ing Point,	°C. 77.	1	100.0
Melting Poin	nt, °C.	-82	.4	0.0
Critical Ten	perature,	°C. 250	.1	374.2
Oritical Pro	ssure, Mm.	Hg 28,8	04	165,680
A	sectropie D	ete ⁸		
Boiling Point. B		mposition, Mole %		sure of Pure ts. Mm. Hg
No. OC.	Mm. Hg Coi	•		Component 2b
1 -1.90	25.0	84.56	21.5	3.986
2 10.05	50.0	83.22	45.1	9.240
2 10.05 3 18.45	78.5	81.99	67.8	15,921
4 19.88	82.2	81.63	71.2	16.873
5 31.35	150.0	78.21	127.2	34.373
6 34.82	176.0	77.51	149.3	41.756
7 42.55	250.0	75.82	209.1	63.295
8 49.06	329.8	73.55	273.1	88.284
9 54.94	420.0	72.38	343.7	117.7
10 56.44	446.2	72.26	264.1	126.4
11 64.33	606.0	70.42	484.4	181.9
12 64.60	613.8	70.22	492.4	184.2
12 69.83	745.0	69.18	592 · O	232.0
14 70.38	760.0	68.85	603.5	237.5
15 74.38	875.0	68.05	692.9	218.6
16 75.28	908.5	67.78	713.0	291.9
17 77.66	984.8	67.38	772.8	322.8
18 82.95	1177.9	66.18	917.6	399.8
19 88.49	1415.0	65.05	1089.7	495.4
20 89.08	1441.3	64.95	1109.5	507.7

aReference: (10).

bReferences: (4,15).

TABLE 82

				Compos	neut 1	Component 2
Sys	tem		Hyd	irogen	Chloride	Water
Formula				HCl		H ₂ O
Formula Weight				36.47		18.016
Normal Boiling Point, OC.				-85		100
Melting Point, OC.				-114.3		0.0
Cri	tical Te	mperatur	e, OC.	51.4		374.2
Cri	tical Pr	essure, l	Vm. Hg	62,0	16	165,680
**************************************	Boiling	sectropic	Data ^a Compositi	- M 23 1	fanor Dro	ssure of Pure
	~	Pressure	Mole %	.VH ₂ 1	Compone	
No.	og.	Mm. Hg	Component	: 1 00		16 Component 20
1	48.724	50	13.126		58,891	86.81
2	69.956		12.557 ^d		-	233.3
3	81.205		12.158		Were affect dates adopt sales with	372.7
4	90.237		11.870°		THE CALL SELECTION AND ADDRESS.	580.51
5	92.080		11.754 ^d 11.655 ^d	•	with the base day the was	568.65
6	95.029	450	2 5 St. 3% Bt. ***			•
					Mildr street Associated below stand	634.58
7	97.578	\$0 0	11.557		7000 1000 1000 1000 1000 1000 4900 1000 1000 1000 1000 1000	634.58 696.55
8	99.658	500 540	11.557 11.471	• •	Might state Ages along bein ages digit state land state was ages micr state ages and along state micr state ages ages ages	624.58 696.55 750.62
8 9	99.653	500 540 500	11.557 11.471 11.385	• • •	### ### ### ### ### ### ### ### ### ###	634.58 696.55 750.62 821.90
8 9 10	99.653 102.209 103.967	\$00 540 600 640	11.557 11.471 11.385 11.305	• · · · · · · · · · · · · · · · · · · ·	This war age are age dip tip tip you wan han me dipi dan ten wan you den wije dan ten wan you ten dip dan ten den ten ten dip dan ten den ten ten dip dan ten den ten ten dip dan ten dan ten dipi dan den den den	634.58 696.55 750.62 821.90 874.06
8 9 10 11	99.653 102.209 103.967 105.564	\$00 540 600 640 680	11.557 11.471 11.385 11.305 11.247		This was now and the new Opp tips you wan has one digit day has the year day top you and day This day now and day	634.58 696.55 750.62 821.90 874.06 923.94
8 9 10 11 12	99.653 102.209 103.967 105.564 106.424	\$00 540 600 640 680 700	11.557 11.471 11.385 11.505 11.247 11.214		THE WAY AND	634.58 696.55 750.62 821.90 674.06 923.94 951.67
8 9 10 11 12 13	99.653 102.209 103.967 105.564 106.424 107.859	\$00 540 600 640 680 700 740	11.557 11.4716 11.385 11.505 11.247 11.2146 11.158		THE THE ROLL AND STON MAN. THE STOP STOP STON MAN. AND THE STOP STON THE THE STON THE STOP STON THE STON THE STON STON THE STON THE STON STON THE ST	634.58 696.55 750.62 821.90 874.06 923.94 951.67 999.59
8 9 10 11 12 13 14	99.653 102.209 103.967 105.564 106.424 107.859 108.584	\$00 540 600 640 680 700 740 760	11.557 11.471 11.385 11.305 11.247 11.214 11.158 11.130		THE THE ASS. AND THE ASS. THE SHE THE THE THE THE THE THE THE THE THE T	634.58 696.55 750.62 821.90 674.06 923.94 951.67 999.59 1024.46
8 9 10 11 12 13	99.653 102.209 103.967 105.564 106.424 107.859	\$00 540 600 640 680 700 740 760	11.557 11.4716 11.385 11.505 11.247 11.2146 11.158		THE WAY AND AND AND AND SOO SOOT THE TOP AND AND THE SOOT THE TOP AND AND AND SOOT THE TOP AND AND AND AND SOOT THE TOP AND AND AND SOOT THE TOP AND AND AND SOOT THE TOP AND AND	624.58 696.55 750.62 821.90 874.06 923.94 951.67 999.59

aReference: (2).

bacference: (20).

CReference: (4,15).

dInterpolated.

TABLE 83

				Compo	nent 1	Component 2
Sys	tem			Hydroge	n Bromide	Water
For	mula			H	Br	и20
Formula Weight				80.92		18.016
Normal Boiling Point, Oc.				-67		100
Mel	ting Po	int, oc.		-8	17.0	0.0
Cri	tical To	emperature	. °C.	90	•0	374.2
Ori	tical P	ressure, i	im. Hg	64,	144	165,680
			. 0			
	Boiling Point,	Azectropic Pressure,	Compos	ition,		sure of Pure
	Boiling		Compos Mole	ition.	Componer	its. Mm. Hg
	Boiling Point,	Pressure,	Compos Mole	ition,	Componer	
No. 1 2	Point,	Pressure,	Compos Kole Compon	ition. % ent 1 91	Component]	its. Mm. Hg
No. 1 2 3	Boiling Point, °C. 74.12	Pressure, Mm. Hg	Compos Mole Compon	ition. % ent 1 91 85	Component]	its. Mm. Hg Lo Component 2 278.52
1 2 3 4	Boiling Point, OC. 74.12 90.35	Pressure. Mm. Hg 100 200	Compos Mole Compon 18.0 17.7	ition. % ent 1 91 85 23	Component]	278.52 532.80 757.56 970.60
1 2 3 4 5	Boiling Point, OC. 74.12 90.35 99.91	Pressure. Mm. Hg 100 200 300	Compos Kole Compon 18.0 17.7 17.5	ition. % ent 1 91 85 20 16	Component]	278.52 532.80 757.56 970.60 1187.81
1 2 3 4 5	74.12 90.35 99.91 107.00 112.94 117.82	Pressure. Mm. Hg 100 200 300 400 500 600	Compos Kole Compon 18.0 17.7 17.5 17.3 17.1	1tion. % ent 1 91 85 23 16 56	Component]	278.52 532.80 757.56 970.60 1187.81 1389.15
1 2 3 4 5 6	74.12 90.35 99.91 107.00 112.94 117.82 122.00	Pressure. Mm. Hg 100 200 200 400 500 600 700	Compos Mole Compon 18.0 17.7 17.5 17.3 17.0 16.9	ition. % ent 1 91 85 28 16 56 20	Component]	278.52 532.80 757.56 970.60 1187.81 1389.15 1586.04
1 2 3 4 5	74.12 90.35 99.91 107.00 112.94 117.82	Pressure. Mm. Hg 100 200 300 400 500 600	Compos Kole Compon 18.0 17.7 17.5 17.3 17.1	ition. % ent 1 91 85 28 16 56 20	Component]	278.52 532.80 757.56 970.60 1187.81 1389.15
No. 1 2 3 4 5 6 7	74.12 90.35 99.91 107.00 112.94 117.82 122.00	Pressure. Mm. Hg 100 200 200 400 500 600 700	Compos Mole Compon 18.0 17.7 17.5 17.3 17.0 16.9	1tion. % ent 1 91 85 22 16 56 20 01	Component]	278.52 532.80 757.56 970.60 1187.81 1389.15 1586.04
1 2 3 4 5 6 7 8	74.12 90.35 99.91 107.00 112.94 117.82 122.00 125.79	Pressure. Mm. Hg 100 200 200 400 500 600 700 800	Compos Mole Compon 18.0 17.7 17.5 17.3 17.1 16.9 16.8 16.7	91 85 23 16 56 20 01 00 10	Component]	278.52 532.80 757.56 970.60 1187.81 1389.15 1586.04 1783.63 1974.01 2158.06
123456789	74.12 90.35 99.91 107.00 112.94 117.82 122.00 125.79 129.13	Pressure. Mm. Hg 100 200 200 400 500 600 700 800 900	Compos Mole Compon 18.0 17.7 17.5 17.3 17.1 17.0 16.9 16.8 16.7	91 85 23 16 56 20 01 00 10	Component]	278.52 532.80 757.56 970.60 1187.81 1389.15 1586.04 1783.63 1974.01

aReference: (1).

bReference: (20).

OReference: (4,15).

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