#### Simple Models For Vapor/Liquid Equilibrium

The preceding section has described what is observed through experimental observation. When thermodynamics is applied to vapour/liquid equilibrium, the goal is to find by calculation the temperatures, pressures, and compositions of phases in equilibrium. Indeed, thermodynamics provides the mathematical framework for the systematic correlation, extension, generalization, evaluation, and interpretation of data. Moreover, it is the means by which the predictions of various theories of molecular physics and statistical mechanics may be applied to practical purposes. None of this can be accomplished without models for the behaviour of systems in vapour/liquid equilibrium. The two simplest are Raoult's law and Henry's law.

### **Raoult's Law**

The two major assumptions required to reduce VLE calculations to Raoult's law are:

- The vapor phase is an ideal gas.
- The liquid phase is an ideal solution.

The first assumption means that Raoult's law can apply only for low to moderate pressures. The second implies that it can have approximate validity only when the species that comprise the system are chemically similar. Just as the ideal gas serves as a standard to which real-gas behaviour may be compared, the ideal solution represents a standard to which real-solution behaviour may be compared. Ideal-solution behaviour is often approximated by liquid phases wherein the molecular species are not too different in size and are of the same chemical nature. Thus, a mixture of isomers, such as ortho-, meta-, and para-xylene, conforms very closely to ideal-solution behaviour. So do mixtures of adjacent members of a homologous series, as for example, n-hexane/n-heptane,

ethanol/propanol, and benzene/toluene. Other examples are acetone/acetonitrile and acetonitrile/nitromethane.

The mathematical expression which reflects the two listed assumptions and which therefore gives quantitative expression to Raoult's law is

$$y_i P = x_i P_i^{\text{sat}}$$
  $(i = 1, 2, ..., N)$  (10.1)

where  $x_i$  is a liquid-phase mole fraction,  $y_i$  is a vapour-phase mole fraction, and  $P_i$  sat is the vapour pressure of pure species i at the temperature of the system. The product  $y_i P$  on the left side of Eq. (10.1) is known as the partial pressure of species i.

A limitation of Raoult's law is that it can be applied only to species for which a vapor pressure is known, and this requires that the species be "subcritical," i.e., that the temperature of application be below the critical temperature of the species.

An important and useful feature of Raoult's law is that it is valid for any species present at a mole fraction approaching unity, provided only that the vapour phase is an ideal gas. Chemical similarity of the constituent species is not here a requirement.

## Dewpoint and Bubblepoint Calculations with Raoult's Law

Although VLE problems with other combinations of variables are possible, engineering interest centres on dewpoint and bubblepoint calculations; there are four classes:

BUBL P: Calculate  $\{y_i\}$  and P, given  $\{x_i\}$  and T DEW P: Calculate  $\{x_i\}$  and P, given  $\{y_i\}$  and T BUBL T: Calculate  $\{y_i\}$  and T, given  $\{x_i\}$  and P DEW T: Calculate  $\{x_i\}$  and T, given  $\{y_i\}$  and P Because  $\Sigma y_i = 1$ , Eq. (10.1) may be summed over all species to yield:

$$P = \sum_{i} x_i P_i^{\text{sat}} \tag{10.2}$$

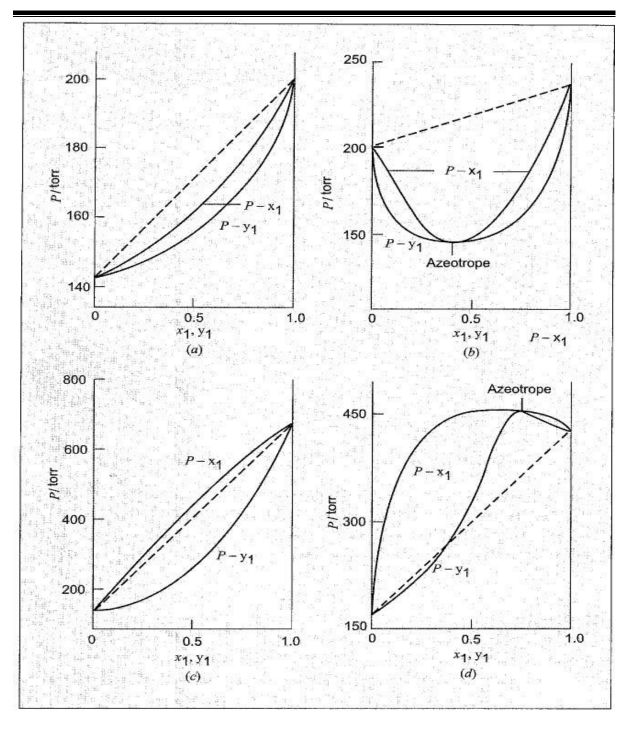
This equation finds application in bubble point calculations, where the vapourphase composition is unknown. For a binary system with  $x_2 = 1 - x_1$ ,

$$P = P_2^{\text{sat}} + (P_1^{\text{sat}} - P_2^{\text{sat}})x_1$$

and a plot of P vs.  $x_1$  at constant temperature is a straight line connecting  $P_2^{sat}$  at  $x_1 = 0$  with  $P_1^{sat}$  at  $x_2 = 1$ . The P-x-y diagrams of Fig. 10.8 show this linear relation. Equation (10.1) may also be solved for xi and summed over all species. With  $\Sigma$  xi = 1, this yields:

$$P = \frac{1}{\sum_{i} y_i / P_i^{\text{sat}}}$$
(10.3)

an equation applied in dewpoint calculations, where liquid-phase compositions are not known.



**Figure 10.8** Pxy diagrams at constant T. (a) Tetrahydrofuran(1)/carbon tetrachloride(2) at 303.15 K (30°C); (b) chloroform(l)/tetrahydrofuran(2) at 303.15 K (30°C) (c) furan(l)/carbon tetrachloride(2) at 303.15 K (30°C); (d) ethanol(l)/toluene(2) at 338.15 K (65°C). **Dashed lines**: Px relation for ideal liquid solutions (Raoult's law)

# Example 10.1

Binary system acetonitrile(1)/nitromethane(2) conforms closely to Raoult's law. Vapor pressures for the pure species are given by the following Antoine equations:

$$\ln P_1^{\text{sat}}/\text{kPa} = 14.2724 - \frac{2,945.47}{t/^{\circ}\text{C} + 224.00}$$
$$\ln P_2^{\text{sat}}/\text{kPa} = 14.2043 - \frac{2,972.64}{t/^{\circ}\text{C} + 209.00}$$

- (a) Prepare a graph showing P vs. x1 and P vs. y1 for a temperature of 75°C.
- (b) Prepare a graph showing t vs. x1 and t vs. y1 for a pressure of 70 kPa.

## Solution 10.1

(a) BUBL P calculations are required. The basis is the binary-system form of Eq. (10.2), repeated here as:

$$P = P_2^{\text{sat}} + (P_1^{\text{sat}} - P_2^{\text{sat}})x_1 \tag{A}$$

At 75°C, by the Antoine equations,

$$P_1^{\text{sat}} = 83.21$$
 and  $P_2^{\text{sat}} = 41.98 \text{ kPa}$ 

Calculations are here very simple, as illustrated for  $x_1 = 0.6$ :

$$P = 41.98 + (83.21 - 41.98)(0.6) = 66.72 \text{ kPa}$$

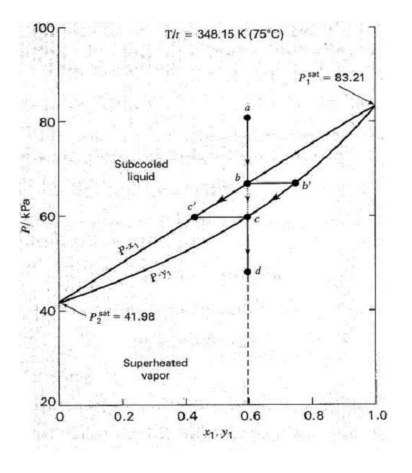
The corresponding value of  $y_1$  is then found from Eq. (10.1):

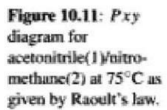
$$y_1 = \frac{x_1 P_1^{\text{sat}}}{P} = \frac{(0.6)(83.21)}{66.72} = 0.7483$$

These results mean that at 75°C a liquid mixture of 60 mol-% acetonitrile and 40 mol-% nitromethane is in equilibrium with a vapor containing 74.83 mol-% acetonitrile at a pressure of 66.72 kPa. The results of calculations for 75°C at a number of values of  $x_1$  are tabulated as follows:

<i>x</i> 1	<i>y</i> 1	P/kPa	xı	yı	P/kPa
0.0	0.0000	41.98	0.6	0.7483	66.72
0.2	0.3313	50.23	0.8	0.8880	74.96
0.4	0.5692	58.47	1.0	1.0000	83.21

These same results are shown by the  $P-x_1-y_1$  diagram of Fig. 10.11. This figure is a phase diagram on which the straight line labeled  $P-x_1$  represents states of saturated liquid; the subcooled-liquid region lies above this line. The curve





labelled P-yi represents states of saturated vapour: the superheated- vapour region lies below this curve points lying between the saturated-liquid and saturated vapour lines are in the two-phase region. What saturated liquid and saturated vapour coexists in equilibrium? The P-x<sub>1</sub> and P-y<sub>1</sub> lines meet at the edges of the diagram what saturated liquid and saturated vapour of the pure species coexist ad the vapour pressures  $P_1^{sat}$  and  $P_2^{sat}$ .

To illustrate the nature of phase behaviour in this binary system we follow the course of a constant-temperature expansion process on the P -xl -y1 diagram, A subcooled liquid mixture of 60 mol-% acetonitrile and 40 mot-% nitmmethane exists in a piston/cylinder arrangement at 75 °C. Its state is represented by point a in Fig. 10.11. Withdrawing the piston slowly enough reduces the pressure while maintaining the system at equilibrium at 75°C. Because the system is closed, the overall composition remains constant during the process and the states of the system as a whole fall on the vertical line descending from point a. When the pressure reaches the value at point b, the system is saturated liquid on the verge of vaporizing. A minuscule further decrease in pressure produces a bubble of vapour, represented by point *b*. The two points b and *b* (x1 =0.6, P = 66.72 kpa. and yi = 0.7483) together represent the state determined by earlier calculations. Point b is a bubble point, and the P-x1 line is the locus of bubble points.

As the pressure is further reduced, the amount of vapour increases and the amount of liquid decreases, with the states of the two phases following paths b'c and b'c respectively. The dotted line from point b to point c represents the overall states of the two-phase system. Finally, as point c is approached, the liquid phase, represented by point c', has almost disappeared, with only droplets (dew) remaining. Point c is therefore a dew point and the P-y1 curve is the locus of dew points. Once the dew has evaporated, only saturated vapour at point c remains, and further pressure reduction leads to superheated vapour at point d.

The composition of the vapour at point c is yi = 0.6, but the composition of the liquid at point c' and the pressure must either be read from the graph or calculated. This is a DEW P calculation, and by Eq. (10.3),

$$P = \frac{1}{y_1/P_1^{\text{saf}} + y_2/P_2^{\text{saf}}}$$

For  $y_1 = 0.6$  and  $t = 75^{\circ}C$ ,

$$P = \frac{1}{0.6/83.21 + 0.4/41.98} = 59.74 \text{ kPa}$$

By Eq. (10.1),

$$x_1 = \frac{y_1 P}{P_1^{\text{sat}}} = \frac{(0.6)(59.74)}{83.21} = 0.4308$$

This is the liquid-phase composition at point c'.

(b) When pressure P is fixed, the temperature varies along with  $x_1$  and  $y_1$ . For a given pressure, the temperature range is bounded by saturation temperatures  $t_1^{sat}$  and  $t_2^{sat}$ , the temperatures at which the pure species exert vapor pressures equal to P. For the present system, these temperatures are calculated from the Antoine equations:

$$t_i^{\text{sat}} = \frac{B_i}{A_i - \ln P} - C_i$$

For P = 70 kPa,  $r_1^{sat} = 69.84^{\circ}$ C and  $t_2^{sat} = 89.58^{\circ}$ C. The simplest way to prepare a *t*-*x*<sub>1</sub>-*y*<sub>1</sub> diagram is to select values of *t* between these two temperatures, calculate  $P_1^{sat}$  and  $P_2^{sat}$  for these temperatures, and evaluate  $x_1$  by Eq. (A), written:

$$x_1 = \frac{P - P_2^{\text{sat}}}{P_1^{\text{sat}} - P_2^{\text{sat}}}$$

For example, at 78°C,  $P_1^{\text{sat}} = 91.76 \text{ kPa}$ ,  $P_2^{\text{sat}} = 46.84 \text{ kPa}$ , and

$$x_1 = \frac{70 - 46.84}{91.76 - 46.84} = 0.5156$$

By Eq. (10.1),

$$y_1 = \frac{x_1 P_1^{\text{sat}}}{P} = \frac{(0.5156)(91.76)}{70} = 0.6759$$

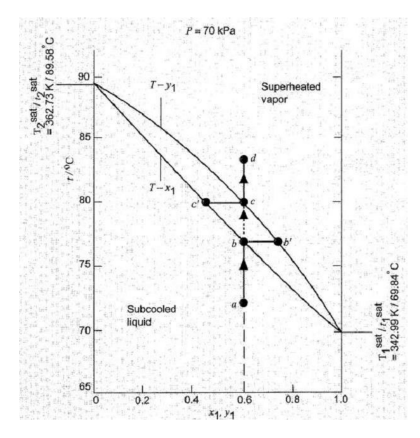
The results of this and similar calculations for P = 70 kPa are as follows:

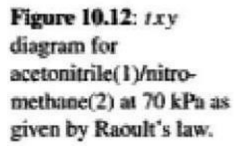
<i>x</i> <sub>1</sub>	<i>y</i> 1	t/°C	xı	<b>y</b> 1	ı∕°C
0.0000	0.0000	89.58 (t2sat)	0.5156	0.6759	78
	0.2401	86	0.7378	0.8484	74
0.3184	0.4742	82	1.0000	1.0000	69.84 (t1sat)

Figure 10.12 is the  $t-x_1-y_1$  diagram showing these results. On this phase diagram, drawn for a constant pressure of 70 kPa, the  $t-y_1$  curve represents states of saturated vapor, with states of superheated vapor lying above it. The  $t-x_1$  curve represents states of saturated liquid, with states of subcooled liquid lying below it. The two-phase region lies between these curves.

With reference to Fig. 10.12, consider a constant-pressure heating process leading from a state of subcooled liquid at point a to a state of superheated vapor at point d. The path shown on the figure is for a constant overall composition of 60 mol-% acetonitrile. The temperature of the liquid increases as the result of heating from point a to point b, where the first bubble of vapor appears. Thus point b is a bubblepoint, and the t- $x_1$  curve is the locus of bubblepoints.

For  $x_1 = 0.6$  and P = 70 kPa, t is determined by a BUBL T calculation, which requires iteration. Equation (10.2) is here written:





$$P_2^{\text{sat}} = \frac{P}{x_1 \alpha + x_2} \tag{B}$$

where  $\alpha = P_1^{\text{sat}}/P_2^{\text{sat}}$ . Subtracting  $\ln P_2^{\text{sat}}$  from  $\ln P_1^{\text{sat}}$  as given by the Antoine equations yields:

$$\ln \alpha = 0.0681 - \frac{2.945.47}{t + 224.00} + \frac{2.972.64}{t + 209.00} \tag{C}$$

The reason for introducing  $\alpha$  is that as the controlling variable it is far less sensitive to *t* than an individual vapor pressure. An initial value of  $\alpha$  is found for an arbitrary intermediate temperature. Iteration is then as follows:

- With the current value of α, calculate P<sub>2</sub><sup>sat</sup> by Eq. (B).
- Calculate t from the Antoine equation for species 2:

$$t = \frac{2,972.64}{14.2043 - \ln P_2^{\text{sat}}} - 209.00$$

- Find a new value of α by Eq. (C).
- · Return to the initial step and iterate to convergence for a final value of t.

The result is  $t = 76.42^{\circ}$ C, the temperature of points b and b'. From the Antoine equation,  $P_1^{sat} = 87.17$  kPa, and by Eq. (10.1) the composition at point b' is:

$$y_1 = \frac{x_1 P_1^{\text{sat}}}{P} = \frac{(0.6)(87.17)}{70} = 0.7472$$

Vaporizing a mixture at constant pressure, unlike vaporizing a pure species, does not in general occur at constant temperature. As the heating process continues beyond point b, the temperature rises, the amount of vapor increases, and the amount of liquid decreases. During this process, the vapor- and liquid-phase compositions change as indicated by paths b'c and bc', until the dewpoint is reached at point c, where the last droplets of liquid disappear. The t- $y_1$  curve is the locus of dewpoints.

The vapor composition at point c is y = 0.6; because the pressure is also known (P = 70 kPa), a *DEW T* calculation is possible. With  $\alpha \equiv P_1^{sat}/P_2^{sat}$ , Eq. (10.3) is written:

$$P_1^{\text{sat}} = P(y_1 + y_2\alpha)$$

The iteration steps are as before, but are based on  $P_1^{\text{sat}}$  rather than  $P_2^{\text{sat}}$ , with

$$t = \frac{2,945.47}{14.2724 - \ln P_1^{\text{sat}}} - 224.00$$

The result here is  $t = 79.58^{\circ}$ C, the temperature of points c and c'. From the Antoine equation,  $P_1^{\text{sat}} = 96.53$  kPa, and Eq. (10.1) gives the composition at point c':

$$x_1 = \frac{y_1 P}{P_1^{\text{sat}}} = \frac{(0.6)(70)}{96.53} = 0.4351$$

Thus the temperature rises from 76.42 to 79.58°C during the vaporization step from point b to point c. Continued heating simply superheats the vapor to point d.