## Solutions

> Mole fraction of component $A=x_{A}$
> Mass Fraction of component $A=m_{A}$
> Volume Fraction of component $A=\phi_{A}$

Typically, we make a binary blend, A + B, with mass fraction, $\mathrm{m}_{\mathrm{A}}$, and want volume fraction, $\phi_{\mathrm{A}}$, or mole fraction, $\mathrm{x}_{\mathrm{A}}$.
$\phi_{A}=\left(m_{A} / \rho_{A}\right) /\left(\left(m_{A} / \rho_{A}\right)+\left(m_{B} / \rho_{B}\right)\right)$
$\mathrm{x}_{\mathrm{A}}=\left(\mathrm{m}_{\mathrm{A}} / \mathrm{MW}_{\mathrm{A}}\right) /\left(\left(\mathrm{m}_{\mathrm{A}} / \mathrm{MW}_{\mathrm{A}}\right)+\left(\mathrm{m}_{\mathrm{B}} / \mathrm{MW}_{\mathrm{B}}\right)\right)$

## Solutions

## $\mathbf{G}=\mathbf{H}-\mathbf{T S} \quad \mathbf{A}=\mathbf{U}-\mathbf{S T} \quad \mathbf{U}=\mathbf{H}-\mathbf{P V}$ Need the Entropy S

Three ways to get entropy and free energy of mixing
A) Isothermal free energy expression, pressure expression
B) Isothermal volume expansion approach, volume expression
C) From statistical thermodynamics

## A. Pressure Expression: Mix two ideal gasses, A and B -S U V

$\mathrm{p}=\mathrm{p}_{\mathrm{A}}+\mathrm{p}_{\mathrm{B}} \quad \mathrm{p}_{\mathrm{A}}$ is the partial pressure $\mathrm{p}_{\mathrm{A}}=\mathrm{x}_{\mathrm{A}} \mathrm{p}$ (Raoult's Ideal Mixing Law) - pGT

For single component molar $G=\mu$
$\mu_{0}$ is at $\mathrm{p}_{0, \mathrm{~A}}=1 \mathrm{bar}$
At pressure $\mathrm{p}_{\mathrm{A}}$ for a pure component isothermal ideal gas
$\mu_{\mathrm{A}}=\mu_{0, \mathrm{~A}}+\mathrm{RT} \operatorname{In}\left(\mathrm{p} / \mathrm{p}_{0, \mathrm{~A}}\right)=\mu_{0, \mathrm{~A}}+\mathrm{RT} \operatorname{In}(\mathrm{p})$
$d G=-S d T+V d p$
Isothermal and Ideal Gas
$\mathrm{dG}=\mathrm{RTdp} / \mathrm{p}$
$\mathrm{G}=\mathrm{RT} \ln \left(\mathrm{p} / \mathrm{p}_{0}\right)$

For a mixture of $A$ and $B$ with a total pressure $p_{\text {tot }}=p_{0, A}=1$ bar and $p_{A}=x_{A} p_{\text {tot }}$
For component A in a binary mixture
$\mu_{A}\left(x_{A}\right)=\mu_{0 . A}+R T \ln \left(x_{A} p_{\text {tot }} / p_{0, A}\right)=\mu_{0 . A}+R T \ln \left(x_{A}\right) \quad$ Isothermal ideal gas (no enthalpy)

Notice that $\mathrm{x}_{\mathrm{A}}$ must be less than or equal to 1 , so $\ln \mathrm{x}_{\mathrm{A}}$ must be negative or 0 So, the chemical potential must drop in the solution for a solution to exist. Ideal gasses only have entropy so entropy drives mixing in this case.

This can be written, $\mathrm{x}_{\mathrm{A}}=\exp \left(\left(\mu_{\mathrm{A}}\left(\mathrm{x}_{\mathrm{A}}\right)-\mu_{0 . \mathrm{A}}\right) / \mathrm{RT}\right)$
Which indicates that $\mathrm{x}_{\mathrm{A}}$ is the Boltzmann probability of finding A

## Mix two real gasses, $A$ and $B$

$$
\begin{gathered}
\mu_{\mathrm{A}}\left(x_{\mathrm{A}}\right)=\mu_{\mathrm{A}}^{\mathrm{o}}+R T \ln \left(\frac{f_{\mathrm{A}}}{p_{\mathrm{A}}^{\mathrm{o}}}\right)=\mu_{\mathrm{A}}^{\mathrm{o}}+R T \ln \left(f_{\mathrm{A}}\right) \quad \text { Gas } \\
\mu_{\mathrm{A}}=\mu_{\mathrm{A}}^{*}+R T \ln a_{\mathrm{A}} \quad \text { Solution } \\
\mu_{\mathrm{A}}^{*}=\mu_{0 . \mathrm{A}} \text { if } \mathrm{p}=1
\end{gathered}
$$

## Solutions

## $\mathbf{G}=\mathbf{H}-\mathbf{T S} \quad \mathbf{A}=\mathbf{U}-\mathbf{S T} \quad \mathbf{U}=\mathbf{H}-\mathbf{P V}$ Need the Entropy S

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## B. Volume Expression: Ideal Gas Mixing

For isothermal $\Delta \mathrm{U}=\mathrm{C}_{\mathrm{V}} \mathrm{dT}=0=\mathrm{dQ}+\mathrm{dW}$
$d Q=-d W=p d V$
For ideal gas $\mathrm{dQ}=-\mathrm{dW}=\mathrm{nRT} \ln \left(\mathrm{V}_{\mathrm{f}} / \mathrm{V}_{\mathrm{i}}\right)$
$d Q=T \Delta S$
$\Delta \mathrm{S}=\mathrm{nR} \ln \left(\mathrm{V}_{\mathrm{f}} / \mathrm{V}_{\mathrm{i}}\right)$
$d U=-p d V+T d S$
Isothermal and Ideal Gas
$\mathrm{dG}=\mathrm{RT} \mathrm{dp} / \mathrm{p}$

Consider a process of expansion of a gas from $V_{A}$ to $V_{\text {tot }}$
The change in entropy is $\Delta \mathrm{S}_{\mathrm{A}}=\mathrm{n}_{\mathrm{A}} \mathrm{R} \ln \left(\mathrm{V}_{\text {tot }} / \mathrm{V}_{\mathrm{A}}\right)=-\mathrm{n}_{\mathrm{A}} \mathrm{R} \ln \left(\mathrm{V}_{\mathrm{A}} / \mathrm{V}_{\text {tot }}\right)$
Consider an isochoric mixing process of ideal gasses A and B .
$A$ is originally in $V_{A}$ and $B$ in $V_{B}$
$V_{\text {tot }}$ is $V_{A}+V_{B}$
The change in entropy for mixing of $A$ and $B$ is
$\Delta \mathrm{S}_{\text {mixing A and } \mathrm{B}}=-\mathrm{n}_{\mathrm{A}} \mathrm{R} \ln \left(\mathrm{V}_{\mathrm{A}} / \mathrm{V}_{\text {tot }}\right)-\mathrm{n}_{\mathrm{B}} \mathrm{R} \ln \left(\mathrm{V}_{\mathrm{B}} / \mathrm{V}_{\text {tot }}\right)=-\mathrm{nR}\left(\mathrm{x}_{\mathrm{A}} \ln \mathrm{x}_{\mathrm{A}}+\mathrm{x}_{\mathrm{B}} \ln \mathrm{x}_{\mathrm{B}}\right)$
For an isothermal, isochoric mixture of ideal gasses (also isobaric since $\mathrm{P} \sim \mathrm{T} / \mathrm{V}$ )
For ideal gasses $\Delta \mathrm{H}_{\text {mixing }}=0$ since there is no interaction
$\Delta \mathrm{G}_{\text {mixing }}=\Delta \mathrm{H}_{\text {mixing }}-\mathrm{T} \Delta \mathrm{S}_{\text {mixing }}=-\mathrm{T} \Delta \mathrm{S}_{\text {mixing }}=\mathrm{nRT}\left(\ln \mathrm{x}_{\mathrm{A}}+\operatorname{lnx}_{\mathrm{B}}\right)$
So, the molar Gibbs Free energy for mixing is $\Delta \mathbf{G}_{\text {mixing }}=\mathbf{R T}\left(\mathbf{x}_{\mathbf{A}} \ln \mathbf{x}_{\mathbf{A}}+\mathbf{x}_{\mathbf{B}} \ln \mathbf{x}_{\mathbf{B}}\right)$

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## $\mathbf{G}=\mathbf{H}-\mathbf{T S} \quad \mathbf{A}=\mathbf{U}-\mathbf{S T} \quad \mathbf{U}=\mathbf{H}-\mathbf{P V}$ Need the Entropy S

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## C. Statistical Thermodynamics

Boltzmann's Law: $\Delta \mathrm{S}=\mathrm{k}_{\mathrm{B}} \ln \Omega$
$\Omega$ Is the number of states
For mixing of $\mathrm{n}_{\mathrm{A}}$ and $\mathrm{n}_{\mathrm{B}}$ with n total molecules
$\Omega=n!/\left(n_{A}!n_{B}!\right)$
Sterling's approximation for large $\mathrm{n}, \mathrm{n}!\sim \mathrm{n} \ln (\mathrm{n})-\mathrm{n}$
We assume that n is large then
$\ln \Omega=-\left(n_{A} \ln \left(n_{A} / n\right)+n_{B} \ln \left(n_{B} / n\right)\right)$
$\Delta S=-k_{B}\left(n_{A} \ln \left(n_{A} / n\right)+n_{B} \ln \left(n_{B} / n\right)\right)=-n k_{B}\left(x_{A} \ln \left(x_{A}\right)+x_{B} \ln \left(x_{B}\right)\right)$
$\Delta G_{\text {mixing }}=\Delta H_{\text {mixing }}-T \Delta S_{\text {mixing }}=-T \Delta S_{\text {mixing }}=n R T\left(x_{A} \ln x_{A}+x_{B} \ln x_{B}\right)$

## Some Types of Entropy

Thermodynamic entropy measured experimentally, Q/T

## Configurational also called Combinatorial

Conformational
Translational and Rotational Entropy (Brownian motion)
Vibrational entropy
Conformational entropy computed in internal or Cartesian coordinates
(which can even be different from each other)
Conformational entropy computed on a lattice
Entropy associated with organization on mixing (Hyrophobic effect and many other unexpected features on mixing)

## An "Ideal Solution" means:

The change on mixing:
$\Delta \mathrm{S}=-\mathrm{nk}_{\mathrm{B}}\left(\mathrm{x}_{\mathrm{A}} \ln \left(\mathrm{x}_{\mathrm{A}}\right)+\mathrm{x}_{\mathrm{B}} \ln \left(\mathrm{x}_{\mathrm{B}}\right)\right)$
Since $(\ln x)$ is always negative or $0, \Delta \mathrm{~S}$ is always positive for ideal solutions
$\Delta \mathrm{G}=-\mathrm{T} \Delta \mathrm{S}$
Since $(\ln x)$ is always negative or $0, \Delta G$ is always negative (or 0 ) and ideal solutions always mix
$\Delta \mathrm{H}$ is 0 , there is no interaction in ideal mixtures, there is no excluded volume, particles are ghosts to each other $\Delta \mathrm{V}=(\mathrm{d} \Delta \mathrm{G} / \mathrm{dp})_{\mathrm{T}}=0$, there is no loss or gain of volume compared to the summed volume

Boltzmann Probability of a Thermally Reversible Event $=\exp (-\Delta G / k T)$
$\Delta G$ is the change in free energy for the event
$\Delta G=\Delta H-T \Delta S$ (Const. $T \& P$, e.g. an experimental measurement)
$\Delta A=\Delta U-T \Delta S$ (Const. $T \& V$, e.g. a simulation on a grid)
Consider an ideal gas with no enthalpic interactions, $\Delta H=0$
The probability of finding an atom A in a mixture of A and B is the molar concentration $x_{\mathrm{A}}$ $\Delta S_{\mathrm{A}} / \mathrm{k}=-\ln x_{\mathrm{A}}$ from the Boltzmann probability
and
$\Delta S / \mathrm{k}=-x_{\mathrm{A}} \ln x_{\mathrm{A}}-x_{\mathrm{B}} \ln x_{\mathrm{B}}$ by a rule of random mixtures
$\Delta G_{\text {ideal }} / \mathrm{k} T=x_{\mathrm{A}} \ln x_{\mathrm{A}}+x_{\mathrm{B}} \ln x_{\mathrm{B}}$ this is always negative since "ln" of a number less than 1 is negative, so mixing always reduces the free energy so it always occurs for an ideal mixture

Hildebrandt Real Solution model considers binary interactions.
The odds of a binary interaction of A and B is $x_{\mathrm{A}} x_{\mathrm{B}}$
This interaction has an average enthalpy (or internal energy) $\Delta H=x_{\mathrm{A}} x_{\mathrm{B}} \Omega_{\mathrm{AB}}$ or $x_{1} x_{2} A_{12}$ Where the first term is the Hildebrandt binary interaction parameter and the second is the Margules one-parameter interaction term. For polymers $\chi_{12}$ is used, the Flory -Huggins interaction parameter (with a few modifications).
$\Delta G_{\text {real solution }} / k T=x_{1} \ln x_{1}+x_{2} \ln x_{2}+x_{1} x_{2} A_{12}=x_{1} \ln a_{1}+x_{2} \ln a_{2}=x_{1} \ln \gamma_{1} x_{1}+x_{2} \ln \gamma_{2} x_{2}$
$\Delta H / \mathrm{k} T=\Delta G_{\text {excess }} / \mathrm{k} T=x_{1} x_{1} A_{12}=x_{1} \ln \gamma_{1}+x_{2} \ln \gamma_{2}$

## Activity Coefficients as Derivatives

$$
G^{E}=R T \sum_{i} x_{i} \ln \left(\gamma_{i}\right) \quad \text { Excess free energy per mole }
$$

$\underline{G}^{\mathrm{E}}$ is the excess free energy value, so $\boldsymbol{n} \boldsymbol{G}^{\mathrm{E}}$
The excess chemical potential of component " i " in phase L or $\mathrm{V}, \mu_{\mathrm{i}}^{\mathrm{E}}$ is given by

$$
\left(\frac{\partial G^{E}}{\partial n_{i}}\right)_{T, P, n_{j+i}}=\bar{G}_{i}^{E}=\mu_{i}^{E}=R T \ln \gamma_{i}
$$

$$
\begin{gather*}
\frac{G^{E}}{R T}=A_{12} x_{1} x_{2} \\
\frac{G^{E}}{R T}=n\left(A_{12} x_{1} x_{2}\right)=\left(A_{12} n_{2}\right)\left(\frac{n_{1}}{n}\right) \\
\frac{1}{R T}\left(\frac{\partial G^{E}}{\partial n_{1}}\right)_{T, P, n_{2}}=\ln \gamma_{1}=A n_{2}\left[\frac{1}{n}-\frac{n_{1}}{n^{2}}\right]=A \frac{n_{2}}{n}\left[1-\frac{n_{1}}{n}\right]=A x_{2}\left(1-x_{1}\right) \\
\ln \gamma_{1}=A x_{2}^{2} ; \text { similarly } \ln \gamma_{2}=A x_{1}^{2}
\end{gather*}
$$

This is how you can find the activity coefficients from the Margules one-parameter coefficient (or the Hildebrandt or Flory-Huggins interaction parameters)

## Real Solutions

## Molar excess functions or departure functions:

Difference between real value and ideal value

$$
\begin{aligned}
& \text { Excess } \Delta G_{\text {mixing }}=\Delta G_{\text {mixing }}-R T\left(x_{A} \ln x_{A}+x_{A} \ln x_{B}\right) \\
& \text { Excess } \Delta S_{\text {mixing }}=\Delta S_{\text {mixing }}+R\left(x_{A} \ln x_{A}+x_{A} \ln x_{B}\right) \\
& \text { Excess } \Delta H_{\text {mixing }}=\Delta H_{\text {mixing }} \\
& \text { Excess } \Delta \mathrm{V}_{\text {mixing }}=\Delta \mathrm{V}_{\text {mixing }}
\end{aligned}
$$



Figure 3.2 Molar volume of mixing of molten $\mathrm{Pb}-\mathrm{Sn}$ at 1040 K [1] and $\mathrm{Pb}-\mathrm{Sb}$ at 907 K [2] as a function of composition.

## Real Solutions

$x_{A}$ becomes $a_{A}$ the activity so

$$
\Delta G_{\text {mixing }}=R T\left(x_{A} \ln a_{A}+x_{A} \ln a_{B}\right)
$$

$$
\text { Excess } \Delta G_{\text {mixing }}=\Delta G_{\text {mixing }}-R T\left(x_{A} \ln x_{A}+x_{B} \ln x_{B}\right)
$$

$$
=R T\left(x_{A} \ln \left(a_{A} / x_{A}\right)+x_{B} \ln \left(a_{B} / x_{B}\right)\right)
$$

$$
=\operatorname{RT}\left(x_{A} \ln \left(\gamma_{A}\right)+x_{B} \ln \left(\gamma_{B}\right)\right)
$$

$\gamma$ Is the activity coefficient

$$
\text { Excess } \Delta S_{\text {mixing }}=-R\left(x_{A} \ln \left(\gamma_{A}\right)+x_{B} \ln \left(\gamma_{B}\right)\right)
$$

Method to use departure functions for calculations (PREOS.xIs)

1) Calculation of properties in the ideal state is simple
2) With an equation of state the departure function can be calculated
3) For any transition first calculate the departure function to the ideal state
4) Then carry out the desired change as an ideal mixture or gas
5) Then use the departure function to return to the real state


Figure 3.5 The molar Gibbs energy of mixing and the molar excess Gibbs energy of mixing of molten $\mathrm{Fe}-\mathrm{Ni}$ at 1850 K . Data are taken from reference [3].

$$
\mathrm{a}=\gamma \mathrm{x} \text { so } \mathrm{da} / \mathrm{dx}=\gamma
$$



Get the activity coefficient at infinite dilution by extrapolating the slope for pure component to $x=0$ This is used for Henry's Law and a few other places.
(Activity coefficient is $a / x$ )

Figure 3.4 The activity of Fe and Ni of molten $\mathrm{Fe}-\mathrm{Ni}$ at 1850 K [3]. At $x_{\mathrm{Ni}}=0.4$ the activity coefficient of Ni is given by MQ/PQ.


Figure 3.4 The activity of Fe and Ni of molten $\mathrm{Fe}-\mathrm{Ni}$ at 1850 K [3]. At $x_{\mathrm{Ni}}=0.4$ the activity coefficient of Ni is given by $\mathrm{MQ} / \mathrm{PQ}$.

We want the infinite dilution activity coefficients to write a function to predict the activity and the activity coefficient.

This is one of the parameters for the functions.

With the function for activity we can predict the free energy and the miscibility.

## Solute

Henry's law for dilute solutions

Vapor pressure of solute $=k_{i} x_{i}$ $\mathrm{k}_{\mathrm{i}}$ is the Henry's Law Constant $\mathrm{x}_{\mathrm{i}}$ is the solute molar fraction (low $\sim 0$ )

## Solvent or ideal mixture

Raoult's Law for solvent or ideal mixtures Vapor pressure $=\mathrm{p}^{*} \mathrm{x}_{\mathrm{j}}$ $\mathrm{x}_{\mathrm{j}}$ is the solvent molar fraction (high $\sim 1$ )
$\mathrm{p}^{*}$ is the vapor pressure of the pure solvent

If a solution is ideal, then $\mathrm{x}_{\mathrm{A}}=\mathrm{a}_{\mathrm{A}}$ and $\gamma_{\mathrm{A}}=1$
At infinite dilution a solvent is ideal (follows Raoult's law) so

But $\gamma_{A}=1$ at $x_{A}=1$, then Raoult's law is followed if
$\left(\mathrm{d} \gamma_{A} / \mathrm{dx}_{\mathrm{A}}\right)_{\mathrm{xA} A>1}=>0$ (See next slide)
A solute follows Henry's Law if
$\left(\mathrm{da}_{\mathrm{B}} / \mathrm{dx}_{\mathrm{B}}\right)_{\mathrm{xB}}=>0=\gamma_{\mathrm{B}}$ infinite dilution

$$
=\left(\mathrm{d} \gamma_{B} x_{B} / d x_{B}\right)_{x B=>0}=\left(\gamma_{B}+x_{B}\left(d \gamma_{B} / d x_{B}\right)\right)_{\times B=>0}
$$

So

$$
\left(x_{B}\left(d \gamma_{B} / d x_{B}\right)\right)_{x B=>0}=0
$$

This isn't that useful


Figure 3.6 (a) The vapour pressure above molten $\mathrm{Si}-\mathrm{Ge}$ at 1723 K [4]. (b) The corresponding activity coefficients of the two components.

Activity of a component in a solution must be defined relative to a standard state, either Henrian (extrapolated) or Raoultian (measured, $\mathrm{x}=1:: \mathrm{a}=1$ )


## You would use Henrian standard state if the solute had limited solubility like PCBs or oil in water and you wanted the infinite dilution of the water component

The activities on the two standard states are related since

$$
\mu_{i}=\mu_{i}^{\mathrm{R}}+R T \ln a_{i}^{\mathrm{R}}=\mu_{i}^{\mathrm{H}}+R T \ln a_{i}^{\mathrm{H}}
$$

which gives

$$
\frac{a_{i}^{\mathrm{R}}}{a_{i}^{\mathrm{H}}}=\exp \left[-\frac{\left(\mu_{i}^{\mathrm{R}}-\mu_{i}^{\mathrm{H}}\right)}{R T}\right]
$$

Figure 3.7 The activity of Ni of molten $\mathrm{Fe}-\mathrm{Ni}$ at 1850 K using both a Raoultian and a Henrian standard state. Data are taken from reference [3].

## Analytic Solution Models

Want to calculate the miscibility of components
Need to generate a phase diagram
Calculation of the vapor pressure
Simplest approach could be a polynomial. However, this generally doesn't inherently have any real meaning. The polynomial just reproduces existing data within experimental limits. Extrapolation is dangerous.

$$
Y\left(x_{\mathrm{B}}\right)=Q_{0}+Q_{1} x_{\mathrm{B}}+Q_{2} x_{\mathrm{B}}^{2}+\ldots+Q_{n} x_{\mathrm{B}}^{n}=\sum_{i=0}^{n} Q_{i} x_{\mathrm{B}}^{i}
$$

Asymmetric feature: vapor pressure
or

$$
Y\left(x_{\mathrm{B}}\right)=x_{\mathrm{A}} x_{\mathrm{B}} \sum_{i=0}^{n} R_{i}\left(x_{\mathrm{A}}-x_{\mathrm{B}}\right)^{i}=x_{\mathrm{B}}\left(1-x_{\mathrm{B}}\right) \sum_{i=0}^{n} R_{i}\left(1-2 x_{\mathrm{B}}\right)^{i}
$$

Symmetric Feature: Phase behavior

$$
\begin{aligned}
\ln \gamma_{\mathrm{B}}= & \ln \gamma_{\mathrm{B}}^{\infty}+\left(\frac{\partial \ln \gamma_{\mathrm{B}}}{\partial x_{\mathrm{B}}}\right)_{x_{\mathrm{B}} \rightarrow 0} x_{\mathrm{B}}+\frac{1}{2}\left(\frac{\partial^{2} \ln \gamma_{\mathrm{B}}}{\partial x_{\mathrm{B}}^{2}}\right)_{x_{\mathrm{B}} \rightarrow 0}
\end{aligned} x_{\mathrm{B}}^{2}+\ldots .
$$

$$
\ln \gamma_{\mathrm{B}}=\ln \gamma_{\mathrm{B}}^{\infty}+\varepsilon_{1}^{\mathrm{B}} x_{\mathrm{B}}+J_{2}^{\mathrm{B}} x_{\mathrm{B}}^{2}
$$

This is similar to a virial expansion


Figure 3.9 An illustration of low-order terms in the Taylor series expansion of $\ln \gamma_{i}$ for dilute solutions using $\ln \gamma_{\mathrm{Tl}}$ for the binary system $\mathrm{Tl}-\mathrm{Hg}$ at 293 K as example. Here $\ln \gamma_{\mathrm{Tl}}^{\infty}=-2.069, \varepsilon_{1}^{\mathrm{Tl}}=10.683$ and $J_{2}^{\mathrm{Tl}}=-14.4$. Data are taken from reference [8].

## Hildebrand Regular Solution Model

The change on mixing:
$\Delta S=-n k_{B}\left(x_{A} \ln \left(x_{A}\right)+x_{B} \ln \left(x_{B}\right)\right)$ Ideal Solution
Since ( $\ln x$ ) is always negative or $0, \Delta S$ is always positive for ideal solutions
$\Delta G=\Delta H-T \Delta S$
Since ( $\ln x$ ) is always negative or $0, \Delta G$ is positive or negative depending on $\Delta H$ :: can mix or demix
Depending on the sign of $\Delta H$
$\Delta V=(d \Delta G / d p)_{T}=0$, there is no loss or gain of volume compared to the summed volume
$\Delta \mathrm{H}=\mathrm{n} \Omega \mathrm{x}_{\mathrm{A}} \mathrm{x}_{\mathrm{B}}$
$\Omega$ is the interaction coefficient or regular solution constant


The equation is symmetric
Figure 3.10 The molar Gibbs energy of mixing of a regular solution A-B for different values of $\Omega / R T$.

## Hildebrand solubility parameter, $\delta$

## $\Delta \mathrm{H}=\mathrm{n} \Omega \mathrm{x}_{\mathrm{A}} \mathrm{x}_{\mathrm{B}}$

$\Omega$ is the interaction coefficient or regular solution constant

Molar Gibbs free energy of mixing
$G_{m}=R T\left(x_{A} \ln \left(x_{A}\right)+x_{B} \ln \left(x_{B}\right)\right)+\Omega x_{A} x_{B}$
$\Omega=\mathrm{zN}_{\mathrm{A}}\left[\mathrm{u}_{\mathrm{AB}}-\left(\mathrm{u}_{\mathrm{AA}}+\mathrm{u}_{\mathrm{BB}}\right) / 2\right]$
$\Delta H_{m}=\frac{\chi \varphi_{A} \varphi_{B} k T}{V_{0}}$ Flory-Huggins chi parameter, $\sim \Omega / k T$

$$
\chi=\frac{V_{0} V_{\mathrm{M}}}{k T}\left(\delta_{\mathrm{T}, \mathrm{~A}}-\delta_{\mathrm{T}, \mathrm{~B}}\right)^{2} \quad \delta_{\mathrm{T}}=\sqrt{\mathrm{CED}}=\sqrt{\frac{E_{\mathrm{coh}}}{V}}
$$

Hansen solubility parameters

$$
\chi=\frac{V_{0} V_{\mathrm{M}}}{k T}\left[\left(\delta_{\mathrm{D}, \mathrm{~A}}-\delta_{\mathrm{D}, \mathrm{~B}}\right)^{2}+\left(\delta_{\mathrm{P}, \mathrm{~A}}-\delta_{\mathrm{P}, \mathrm{~B}}\right)^{2}+\left(\delta_{\mathrm{H}, \mathrm{~A}}-\delta_{\mathrm{H}, \mathrm{~B}}\right)^{2}\right] \quad \delta_{\mathrm{D}}=\sqrt{\frac{E_{\mathrm{coh}, \mathrm{D}}}{V}}, \delta_{\mathrm{P}}=\sqrt{\frac{E_{\mathrm{coh}, \mathrm{P}}}{V}}, \delta_{\mathrm{H}}=\sqrt{\frac{E_{\mathrm{coh}, \mathrm{H}}}{V}}
$$

## Hildebrand Solubility Parameter, $\delta$

Two materials with similar $\delta$ are miscible
by Hildebrand and Hansen. ${ }^{15,16}$ The Hildebrand solubility parameter $\left(\delta_{\mathrm{T}}\right)$ is the square root of cohesive energy density (CED), and the CED is simply the cohesive energy ( $E_{\text {coh }}$ ) per unit of volume $(V))^{1}$

$$
\begin{equation*}
\delta_{\mathrm{T}}=\sqrt{\mathrm{CED}}=\sqrt{\frac{E_{\mathrm{coh}}}{V}} \tag{1}
\end{equation*}
$$

The intermolecular interactions are composed of dispersive ( $E_{\text {coh }, \mathrm{D}}$ ), polar $\left(E_{\text {cobh, } \mathrm{P}}\right)$, and hydrogen bonding ( $E_{\mathrm{coh}, \mathrm{H}}$ ) interactions, so the Hansen solubility parameters $\left(\delta_{\mathrm{i}}, \mathrm{i}=\mathrm{D}\right.$, $\mathrm{P}, \mathrm{H})$ were proposed as follows: ${ }^{18}$

$$
\begin{equation*}
\delta_{\mathrm{D}}=\sqrt{\frac{E_{\mathrm{coh}, \mathrm{D}}}{V}}, \quad \delta_{\mathrm{p}}=\sqrt{\frac{E_{\mathrm{coh}, \mathrm{P}}}{V}}, \quad \delta_{\mathrm{H}}=\sqrt{\frac{E_{\mathrm{coh}, \mathrm{H}}}{V}} \tag{2}
\end{equation*}
$$

Then the relation between Hansen and Hildebrand solubility parameters as follows:

$$
\begin{equation*}
\delta_{\mathrm{T}}^{2}=\delta_{\mathrm{D}}^{2}+\delta_{\mathrm{P}}^{2}+\delta_{\mathrm{H}}^{2} \tag{3}
\end{equation*}
$$

London dispersion forces (attractive)

Luo Y, Chen X, Wu S, Ca S Luo Z, Shi Y, Molecular Dynamics Simulation Study on Two-Component Solubility Parameters of Carbon Nanotubes and Precisely Tailoring the Thermodynamic Compatibility between Carbon Nanotubes and Polymers, Langmuir 36 9291-9305 (2020).

$$
\Delta G_{\mathrm{m}}=\Delta H_{\mathrm{m}}-T \Delta S_{\mathrm{m}}
$$

$$
\begin{gathered}
\Delta H_{m}=\frac{\chi \varphi_{A} \varphi_{\mathrm{B}} k T}{V_{0}} \quad \begin{array}{l}
\text { Flory-Huggins Equation } \\
\chi^{\sim} 1 / \mathrm{T}
\end{array} \\
\Delta H_{\mathrm{m}}=\varphi_{\mathrm{A}} \varphi_{\mathrm{B}}\left(\delta_{\mathrm{T}, \mathrm{~A}}-\delta_{\mathrm{T}, \mathrm{~B}}\right)^{2} V_{\mathrm{M}}
\end{gathered}
$$

Hildebrand and Scratchard

$$
\chi=\frac{V_{0} V_{\mathrm{M}}}{k T}\left(\delta_{\mathrm{T}, \mathrm{~A}}-\delta_{\mathrm{T}, \mathrm{~B}}\right)^{2}
$$

$$
\chi=\frac{V_{0} V_{\mathrm{M}}}{k T}\left[\left(\delta_{\mathrm{D}, \mathrm{~A}}-\delta_{\mathrm{D}, \mathrm{~B}}\right)^{2}+\left(\delta_{\mathrm{P}, \mathrm{~A}}-\delta_{\mathrm{P}, \mathrm{~B}}\right)^{2}+\left(\delta_{\mathrm{H}, \mathrm{~A}}-\delta_{\mathrm{H}, \mathrm{~B}}\right)^{2}\right]
$$

## Asymmetric equations for asymmetric phase diagram

Sub-regular solution model

$$
\begin{aligned}
\Delta_{\mathrm{mix}}^{\operatorname{exc}} G_{\mathrm{m}} & =x_{\mathrm{A}} x_{\mathrm{B}}\left(A_{21} x_{\mathrm{A}}+A_{12} x_{\mathrm{B}}\right) \\
& \quad \ln \gamma_{1}=x_{2}^{2}\left[A_{12}+2\left(A_{21}-A_{12}\right) x_{1}\right] ; \text { similarly } \ln \gamma_{2}=x_{1}^{2}\left[A_{21}+2\left(A_{12}-A_{21}\right) x_{2}\right] \\
& A_{12}=\left(2-\frac{1}{x_{2}}\right) \frac{\ln \gamma_{1}}{x_{2}}+\frac{2 \ln \gamma_{2}}{x_{1}} \quad A_{21}=\left(2-\frac{1}{x_{1}}\right) \frac{\ln \gamma_{2}}{x_{1}}+\frac{2 \ln \gamma_{1}}{x_{2}} \\
\Delta_{\mathrm{mix}}^{\operatorname{exc}} G_{\mathrm{m}} & =\sum_{i=1}^{m} \sum_{j=1}^{n} x_{\mathrm{A}}^{i} x_{\mathrm{B}}^{j} A_{i j}
\end{aligned}
$$

Redlich-Kister Expression

$$
\begin{aligned}
\Delta_{\operatorname{mix}}^{\mathrm{exc}} G_{\mathrm{m}}= & x_{\mathrm{A}} x_{\mathrm{B}}\left[\Omega+A_{1}\left(x_{\mathrm{A}}-x_{\mathrm{B}}\right)+A_{2}\left(x_{\mathrm{A}}-x_{\mathrm{B}}\right)^{2}\right. \\
& \left.+A_{3}\left(x_{\mathrm{A}}-x_{\mathrm{B}}\right)^{3}+\ldots\right]
\end{aligned}
$$



Figure 11.3. Illustration of calculation of $\mathbf{G}^{\mathbf{E}}$ from experiment and fitting of Margules models to a single point as discussed in Examples 11.1 and 11.5, for 2-propanol + water, with the experimental data points from Fig. 10.8 on page 395. Data are tabulated in Example 11.8. The van Laar model fit to a single point is explained in Section 12.2.


Figure 11.5. (a) One-parameter and two-parameter Margules equation fitted to a single measurement in Examples 11.2 and 11.5 compared with the experimental data points from Fig. 10.8 on page 395. Data are tabulated in Example 11.8. (b) Activity coefficients predicted from the parameters fitted in Example 11.5 compared with points calculated from the data.

$$
\begin{gathered}
\begin{array}{c}
\text { Non-ideal entropy of mixing } \\
\text { Quasi-regular solution model }
\end{array} \\
\Delta_{\mathrm{mix}}^{\mathrm{exc}} G_{\mathrm{m}}=x_{\mathrm{A}} x_{\mathrm{B}} \Omega\left(1-\frac{T}{\tau}\right) \\
\Delta_{\operatorname{mix}}^{\operatorname{exc}} S_{\mathrm{m}}=-\frac{\partial\left(\Delta_{\operatorname{mix}}^{\operatorname{exc}} G_{\mathrm{m}}\right)}{\partial T}=x_{\mathrm{A}} x_{\mathrm{B}}\left(\frac{\Omega}{\tau}\right) \quad \begin{array}{l}
\text { This is a non- } \\
\text { combinatorial } \\
\text { entropy }
\end{array}
\end{gathered}
$$

This would occur if there were ordering on mixing, say when you add oil to water. The enthalpy of mixing favors mixing!!! Water organizes at the surface of oil to a great extent; this reduces entropy and makes oil and water demix. Water actually "likes" oil. This is called the "hydrophobic effect" and is an important concept for protein folding.

## Graphical Method to Estimate Chemical Potential



Figure 3.12 The integral molar Gibbs energy of liquid $\mathrm{Ge}-\mathrm{Si}$ at 1500 K with pure liquid Ge and solid Si as standard states. Data are taken from reference [4].

## Solving the Regular Solution Model

We know that the excess Gibbs free energy for mixing is given by:
$\mathrm{G}^{\mathrm{E}} / \mathrm{RT}=\mathrm{x}_{\mathrm{A}} \ln \gamma_{\mathrm{A}}+\mathrm{x}_{\mathrm{B}} \ln \gamma_{\mathrm{B}} \quad$ Generic expression using activity coefficient
And we have defined for the regular solution model that
$\mathrm{G}^{\mathrm{E}} / \mathrm{RT}=\Omega \mathrm{x}_{\mathrm{A}} \mathrm{x}_{\mathrm{B}}$ Hildebrand Regular Solution expression
If we propose the answer RT $\ln \gamma_{A}=\Omega \mathrm{x}_{\mathrm{B}}{ }^{2}$
We find by substitution generic expression that it equals the regular solution expression since $\mathrm{x}_{\mathrm{B}}+\mathrm{x}_{\mathrm{A}}=1$ (Solving this directly see next slide.)

$$
\begin{aligned}
& \frac{\Delta_{\operatorname{mix}}^{\operatorname{exc}} \bar{G}_{\mathrm{A}}}{R T}=\ln \gamma_{\mathrm{A}}=\frac{\Omega}{R T} x_{\mathrm{B}}^{2} \\
& \frac{\Delta_{\operatorname{mix}}^{\mathrm{exc}} \bar{G}_{\mathrm{B}}}{R T}=\ln \gamma_{\mathrm{B}}=\frac{\Omega}{R T} x_{\mathrm{A}}^{2}
\end{aligned}
$$

$$
G^{E}=R T \sum_{i} x_{i} \ln \left(y_{i}\right)
$$

## Activity Coefficients as Derivatives

$$
\left(\frac{\partial \underline{G}^{E}}{\partial n_{i}}\right)_{T, P, n_{j \neq i}}=\bar{G}_{i}^{E}=\mu_{i}^{E}=R T \ln \gamma_{i}
$$

11.28
( Aclivily coelifcients are related to the partial molar excess Gibbs energy-

$$
\begin{align*}
& \frac{G^{E}}{R T}=A_{12} x_{1} x_{2} \\
& \frac{G^{E}}{R T}=n\left(A_{12} x_{1} x_{2}\right)=\left(A_{12} n_{2}\right)\left(\frac{n_{1}}{n}\right) \\
& \frac{1}{R T}\left(\frac{\partial \underline{\underline{G}}^{E}}{\partial n_{1}}\right)_{T, P, n_{2}}=\ln \gamma_{1}=A n_{2}\left[\frac{1}{n}-\frac{n_{1}}{n^{2}}\right]=A \frac{n_{2}}{n}\left[1-\frac{n_{1}}{n}\right]=A x_{2}\left(1-x_{1}\right) \quad 11.30
\end{align*}
$$

$$
\ln \gamma_{1}=A x_{2}^{2} ; \text { similarly } \ln \gamma_{2}=A x_{1}^{2}
$$

Use of the Gibbs-Duhem Equation to determine the activity of a component

$$
\begin{aligned}
& n_{\mathrm{A}} \mathrm{~d} \mu_{\mathrm{A}}+n_{\mathrm{B}} \mathrm{~d} \mu_{\mathrm{B}}=0 \quad \text { Constant } \mathrm{p}, \mathrm{~T} \\
& x_{\mathrm{A}} \mathrm{~d} \ln a_{\mathrm{A}}+x_{\mathrm{B}} \mathrm{~d} \ln a_{\mathrm{B}}=0 \\
& x_{\mathrm{A}} \mathrm{~d} \ln x_{\mathrm{A}}+x_{\mathrm{A}} \mathrm{~d} \ln \gamma_{\mathrm{A}}+x_{\mathrm{B}} \mathrm{~d} \ln x_{\mathrm{B}}+x_{\mathrm{B}} \mathrm{~d} \ln \gamma_{\mathrm{B}}=0 \\
& x_{\mathrm{A}} \mathrm{~d} \ln x_{\mathrm{A}}+x_{\mathrm{B}} \mathrm{~d} \ln x_{\mathrm{B}}=x_{\mathrm{A}} \frac{\mathrm{~d} x_{\mathrm{A}}}{x_{\mathrm{A}}}+x_{\mathrm{B}} \frac{\mathrm{~d} x_{\mathrm{B}}}{x_{\mathrm{B}}}=\mathrm{d} x_{\mathrm{A}}+\mathrm{d} x_{\mathrm{B}}=0 \\
& x_{\mathrm{A}} \mathrm{~d} \ln \gamma_{\mathrm{A}}+x_{\mathrm{B}} \mathrm{~d} \ln \gamma_{\mathrm{B}}=0 \quad \text { Restatement of Gibbs-Duhem for Solutions } \\
& \quad \text { If you know } \gamma_{\mathrm{A}} \text { you can obtain } \gamma_{\mathrm{B}} \text { by integration } \\
& \ln \gamma_{\mathrm{B}}-\ln \gamma_{\mathrm{B}}\left(x_{\mathrm{B}}=1\right)=-\int_{x_{\mathrm{B}}=1}^{x_{\mathrm{B}}} \frac{x_{\mathrm{A}}}{x_{\mathrm{B}}} \mathrm{~d} \ln \gamma_{\mathrm{A}}
\end{aligned}
$$



Figure 3.13 $x_{\mathrm{Ni}} / x_{\mathrm{Fe}}$ versus $\ln \gamma_{\mathrm{Ni}}$ of molten $\mathrm{Fe}-\mathrm{Ni}$ at 1850 K . Data are taken from reference [3].

