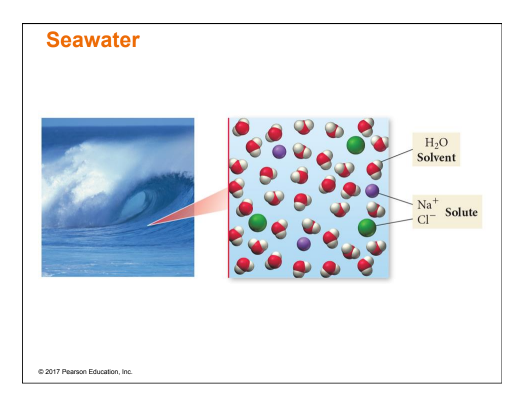


Thirsty Seawater

- Drinking seawater can cause dehydration.
- Seawater
 - is a homogeneous mixture of salts with water.
 - contains higher concentrations of salts than the salt content of your cells.
- As seawater passes through your body, it pulls water out of your cells, due mainly to nature's tendency toward spontaneous mixing.
- This reduces your cells' water level and usually results in diarrhea as this extra liquid flows out with the seawater.



Solutions

- The majority component of a solution is called the **solvent**.
- The minority component is called the **solute**.
- Solutions form in part because of intermolecular forces.
 - The particles of the solute interact with the particles of the solvent through intermolecular forces.

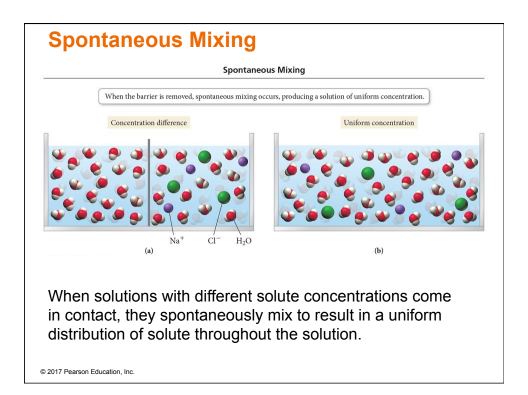
Seawater

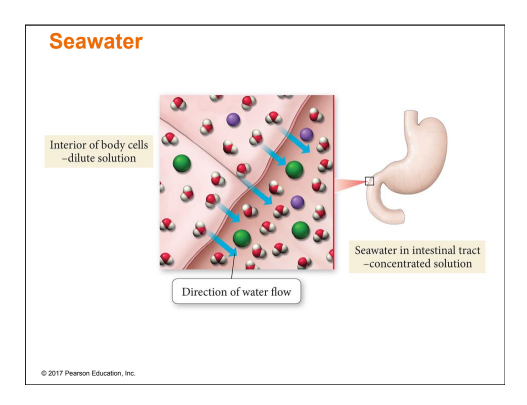
- Drinking seawater will dehydrate you and give you diarrhea.
- The cell wall acts as a barrier to the solute moving, so the only way for the seawater and the cell solution to have uniform mixing is for water to flow out of the cells of your intestine and into your digestive tract.

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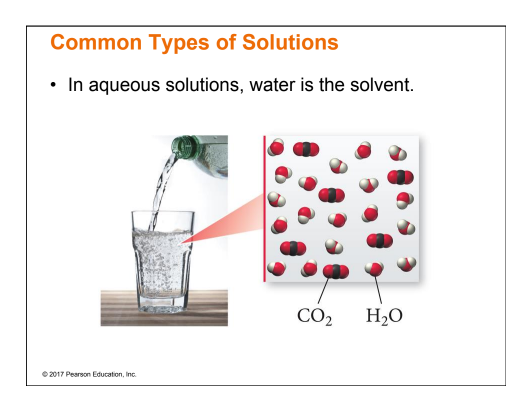
Homogeneous Mixtures

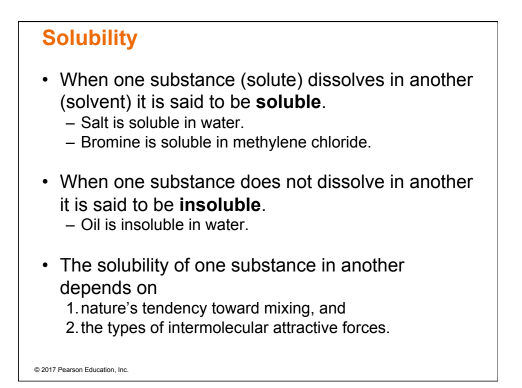
- · A mixture of two or more substances
- Composition may vary from one sample to another
- Appears to be one substance, though really contains multiple materials
- Most homogeneous materials we encounter are actually solutions.
 - For example, air and seawater
- Nature has a tendency toward spontaneous mixing.
 - Generally, uniform mixing is more energetically favorable.

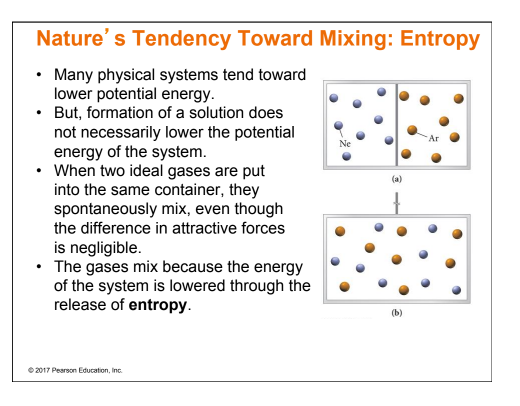


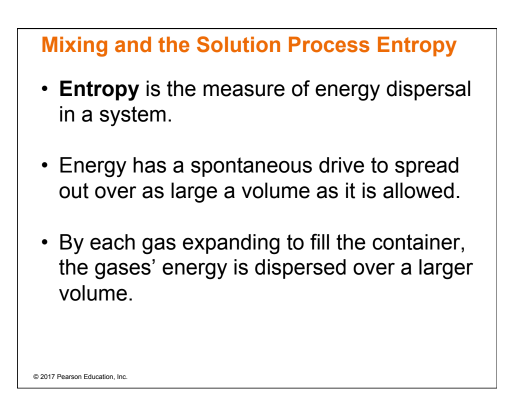


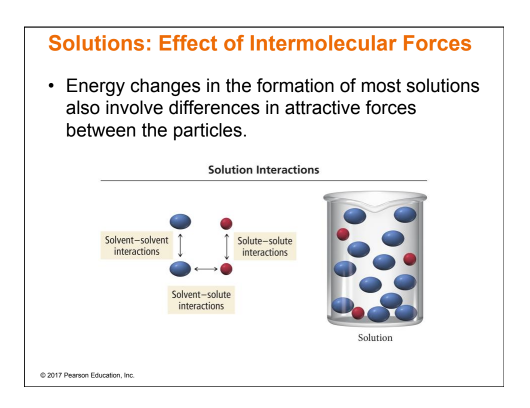
	n may be o her combi	composed nation.	ns of a solid and a liquid
Solution Phase	Solute Phase	Solvent Phase	Example
Gaseous solution	Gas	Gas	Air (mainly oxygen and nitrogen)
Liquid solution	Gas	Liquid	Club soda (CO ₂ and water)
	Liquid	Liquid	Vodka (ethanol and water)
	Solid	Liquid	Seawater (salt and water)
Solid solution	Solid	Solid	Brass (copper and zinc) and other alloys
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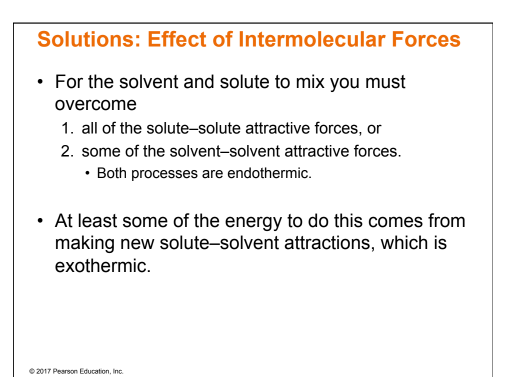


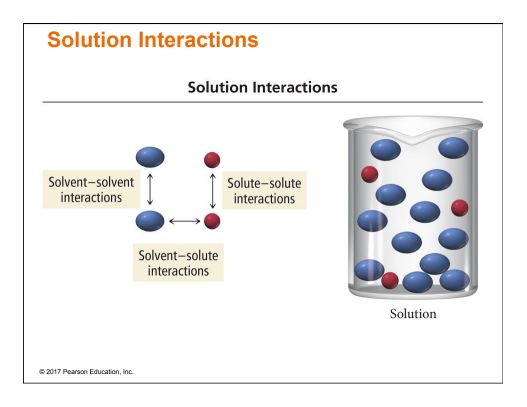


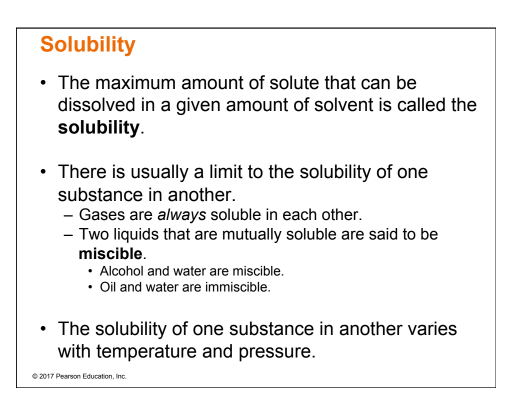


Relative Formati		eractions and	Solution
TABLE 13.2	Relativ	e Interactions and Solution	Formation
Solvent–solute interactions	>	Solvent–solvent and solute–solute interactions	Solution forms
Solvent–solute interactions	=	Solvent–solvent and solute–solute interactions	Solution forms
Solvent-solute	<	Solvent-solvent and solute-solute interactions	Solution may or may not form, depending on relative disparity

 When the solute-to-solvent attractions are weaker than the sum of the solute-to-solute and solventto-solvent attractions, the solution will form only if the energy difference is small enough to be overcome by the increase in entropy from mixing.







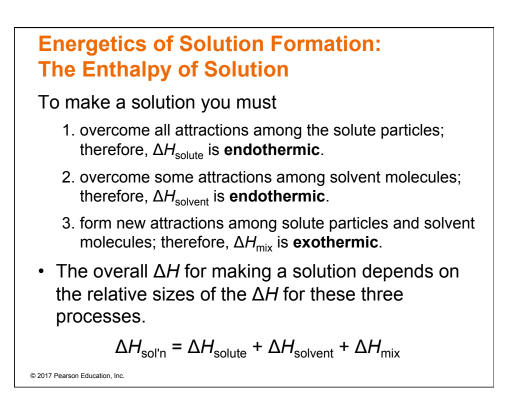
Will It Dissolve?

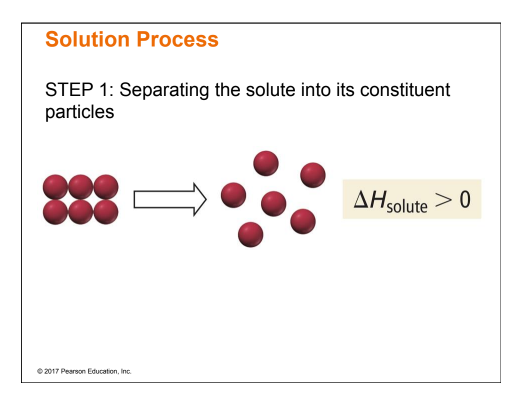
- Chemist's rule of thumb: like dissolves like.
- A chemical will dissolve in a solvent if it has a similar structure to the solvent.
- Polar molecules and ionic compounds will be more soluble in polar solvents.
- Nonpolar molecules will be more soluble in nonpolar solvents.

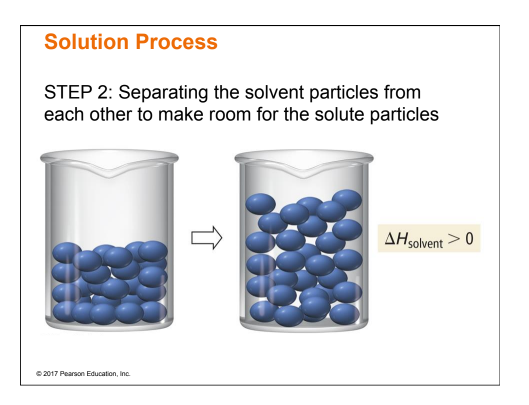
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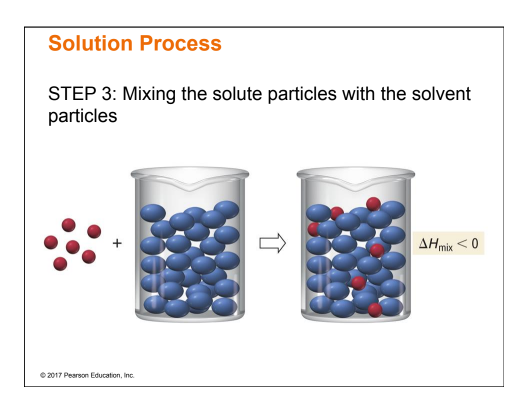
Heat of Solution

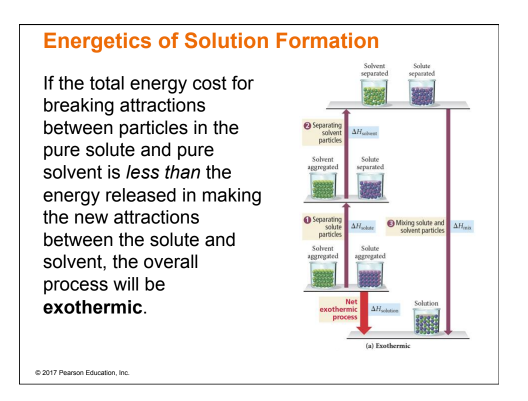
- When some compounds, such as NaOH, dissolve in water, a lot of heat is released.
 - The container gets hot.
- When other compounds, such as NH₄NO₃, dissolve in water, heat is absorbed from the surroundings.
 - The container gets cold.
- Why is this?







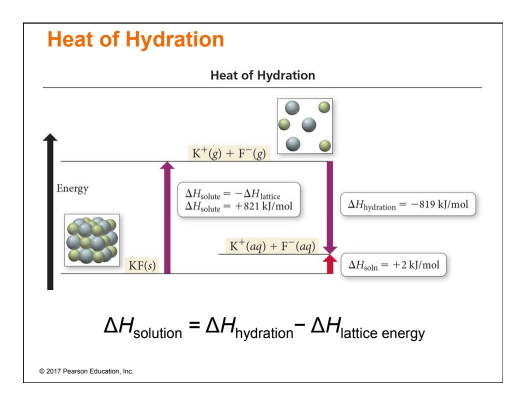


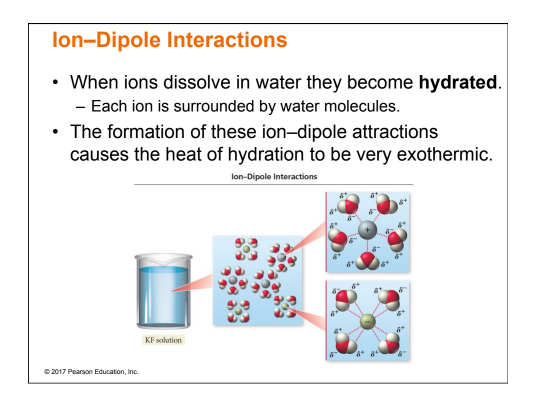


Energetics of Solution Formation Solute If the total energy cost for separated breaking attractions between particles in the Separating $\Delta H_{solvent}$ solvent pure solute and pure particle Solvent Solute solvent is greater than the aggregat epara energy released in making the new attractions Separating Mixing solute and solvent particles ΔH_{colute} $\Delta H_{\rm mix}$ solute between the solute and solvent, the overall Solution process will be Solvent Solut endothermic. aggregated aggregat dothermi (b) Endothermic



- For aqueous solutions of ionic compounds, the energy added to overcome the attractions between water molecules and the energy released in forming attractions between the water molecules and ions is combined into a term called the **heat of** hydration.
 - Attractive forces between ions = lattice energy
 - $\Delta H_{\text{solute}} = -\Delta H_{\text{lattice energy}}$
 - Attractive forces in water = H bonds
 - Attractive forces between ion and water = ion-dipole
 - $-\Delta H_{\text{hydration}}$ = heat released when one mole of gaseous ions dissolves in water = $\Delta H_{\text{solvent}}$ + ΔH_{mix}





Heats of Solution for Ionic Compounds

• For an aqueous solution of an ionic compound, the $\Delta H_{\text{solution}}$ is the difference between the heat of hydration and the lattice energy.

$$\Delta H_{\text{solution}} = \Delta H_{\text{solute}} + \Delta H_{\text{solvent}} + \Delta H_{\text{mix}}$$

$$\Delta H_{\text{solution}} = -\Delta H_{\text{lattice energy}} + \Delta H_{\text{solvent}} + \Delta H_{\text{mix}}$$

$$\Delta H_{\text{solution}} = \Delta H_{\text{hydration}} - \Delta H_{\text{lattice energy}}$$

$$\Delta H_{\text{solution}} = -\Delta H_{\text{lattice energy}} + \Delta H_{\text{hydration}}$$
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- Because the lattice energy is always exothermic, the size and sign on the $\Delta H_{sol'n}$ tells us something about $\Delta H_{hvdration}$.
- If the heat of solution is large and exothermic, then the amount of energy it costs to separate the ions is less than the energy released from hydrating the ions.

 $\Delta H_{\text{hydration}} > \Delta H_{\text{solute}}$ when $\Delta H_{\text{sol'n}}$ is (-).

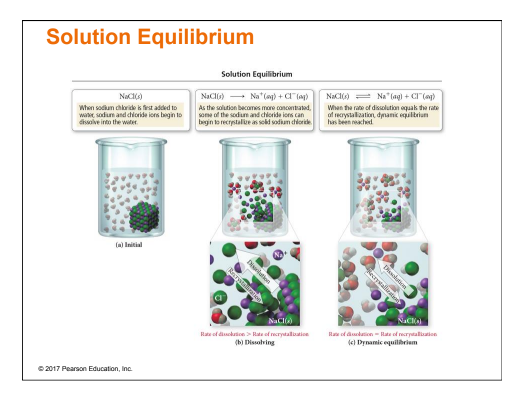
• If the heat of solution is large and endothermic, then the amount of energy it costs to separate the ions is more than the energy released from hydrating the ions.

 $\Delta H_{\text{hydration}} < \Delta H_{\text{solute}}$ when $\Delta H_{\text{sol'n}}$ is (+).

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Solution Equilibrium

- The dissolution of a solute in a solvent is an equilibrium process.
- Initially, when there is no dissolved solute, the only process possible is dissolution.
- Shortly after some solute is dissolved, solute particles can start to recombine to reform solute molecules, but when the rate of dissolution >> the rate of deposition, the solute continues to dissolve.
- Eventually, the rate of dissolution = the rate of deposition—the solution is saturated with solute, and no more solute will dissolve.



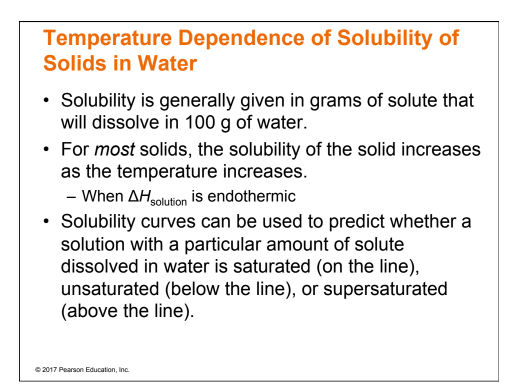
Solubility Limit

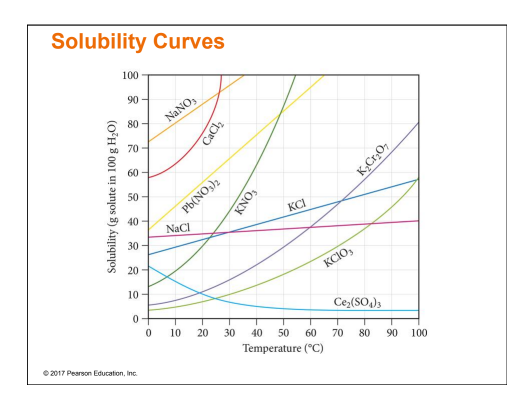
- A solution that has the solute and solvent in dynamic equilibrium is said to be **saturated**.
 - If you add more solute it will not dissolve.
 - The saturation concentration depends on the temperature and pressure of gases.
- A solution that has less solute than saturation is said to be **unsaturated**.
 - More solute will dissolve at this temperature.
- A solution that has more solute than saturation is said to be **supersaturated**.

How Can You Make a Solvent Hold More Solute?

- Solutions can be made saturated at non-room conditions and then can be allowed to come to room conditions slowly.
- For some solutes, instead of coming out of solution when the conditions change, they get stuck between the solvent molecules, and the solution becomes supersaturated.
- Supersaturated solutions are unstable and lose all the solute above saturation when disturbed.
 - For example, shaking a carbonated beverage







Purification by Recrystallization

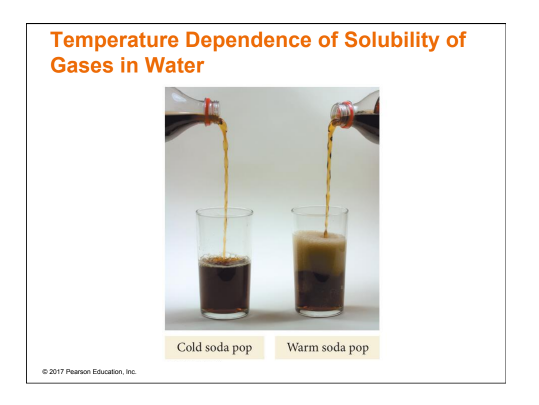
- One of the common operations performed by a chemist is removing impurities from a solid compound.
- One method of purification involves dissolving a solid in a hot solvent until the solution is saturated.
- As the solution slowly cools, the solid crystallizes out, leaving impurities behind.

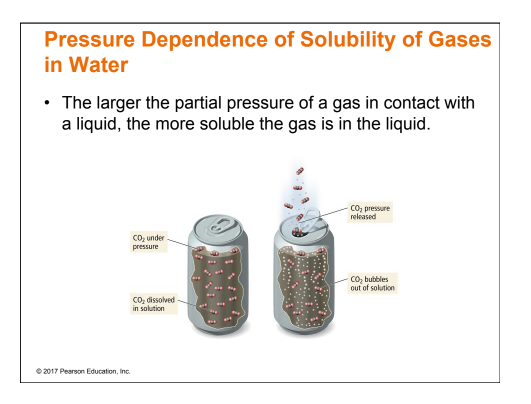


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Temperature Dependence of Solubility of Gases in Water

- Gases generally have lower solubility in water than ionic or polar covalent solids because most are nonpolar molecules.
 - Gases with high solubility usually are actually reacting with water.
- For *all* gases, the solubility of the gas decreases as the temperature increases.
 - The $\Delta H_{\text{solution}}$ is exothermic because you do not need to overcome solute–solute attractions.





Henry's Law

• The solubility of a gas (S_{gas}) is directly proportional to its partial pressure, (P_{gas}) .

$$S_{\rm gas} = k_{\rm H} P_{\rm gas}$$

*k*_H is called the Henry's law constant.

	3.4 Henry's Law for Several Gases at 25 °C
Gas	<i>k</i> _н (M/atm)
02	$1.3 imes 10^{-3}$
N ₂	$6.1 imes 10^{-4}$
CO ₂	$3.4 imes 10^{-2}$
NH ₃	$5.8 imes10^1$
He	$3.7 imes10^{-4}$

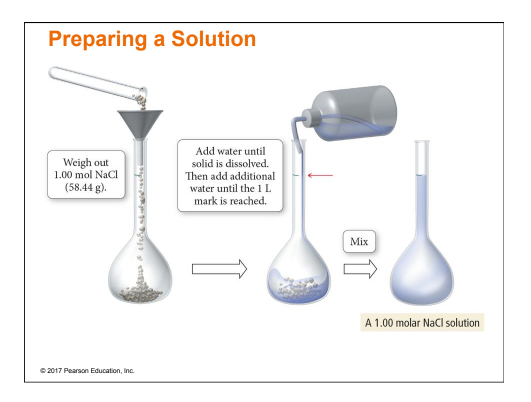
Concentrations

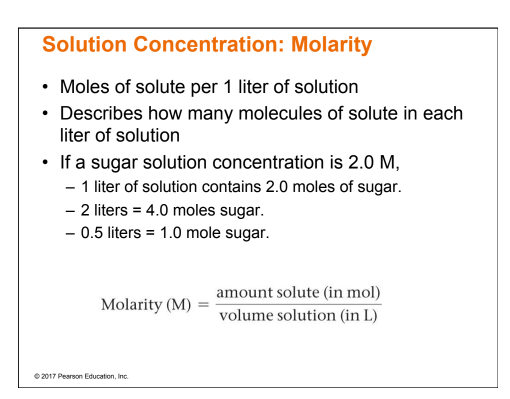
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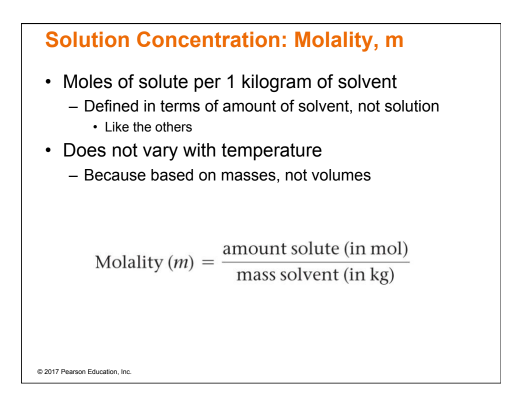
- Solutions have variable composition.
- To describe a solution, you need to describe the components *and* their relative amounts.
- The terms **dilute** and **concentrated** can be used as qualitative descriptions of the amount of solute in solution.
- **Concentration** = amount of solute in a given amount of solution.
 - Occasionally per amount of solvent

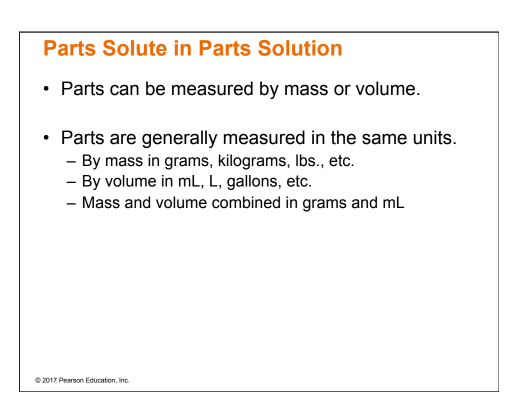
TABLE 13.5 Solution Concentration Terms			
Unit	Definition	Units	
Molarity (M)	amount solute (in mol)	mol	
	volume solution (in L)	L	
Molality (m)	amount solute (in mol)		
	mass solvent (in kg)	kg	
Mole fraction (χ)	amount solute (in mol)		
	total amount of solute and solvent (in mol)		
Mole percent (mol %)	amount solute (in mol)		
	1000000000000000000000000000000000000		
Parts by mass	$\frac{\text{mass solute}}{\text{mass solution}} \times \frac{\text{multiplication factor}}{\text{multiplication factor}}$		
Percent by mass (%)	Multiplication factor = 100	%	
Parts per million by mass (ppm)	Multiplication factor = 10^6	ppm	
Parts per billion by mass (ppb)	Multiplication factor = 10^9	ppb	
Parts by volume (%, ppm, ppb)	$\frac{\text{volume solute}}{\text{volume solution}} \times \text{multiplication factor*}$		

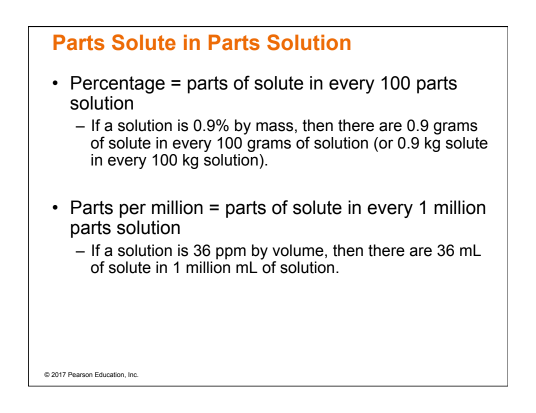
Preparing a Solution Need to know amount of solution and concentration of solution. Calculate the mass of solute needed. Start with an amount of solution. Use concentration as a conversion factor. 5% by mass ⇒ 5 g solute ≡ 100 g solution "Dissolve the grams of solute in enough solvent to total the total amount of solution."

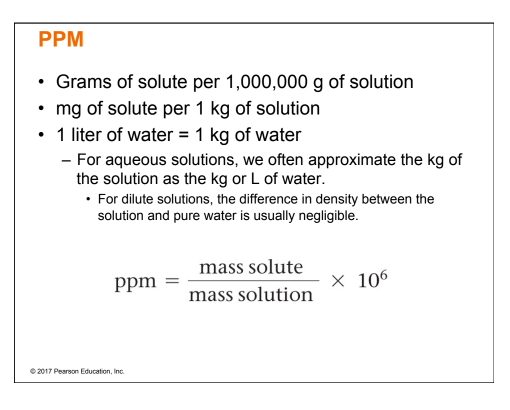






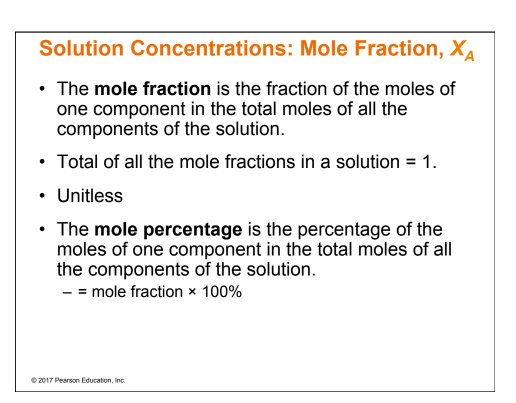






Parts per Billion Concentration

$$ppb = \frac{mass \ solute}{mass \ solution} \times 10^9$$



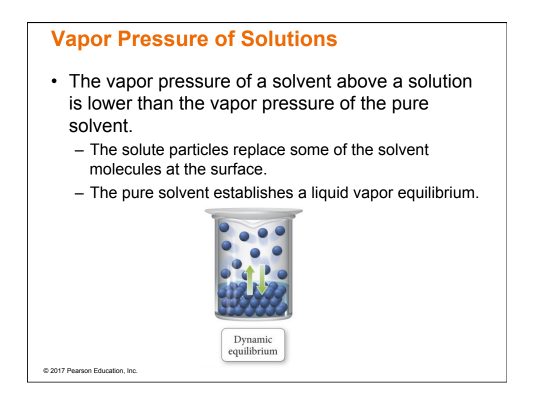
Converting Concentration Units

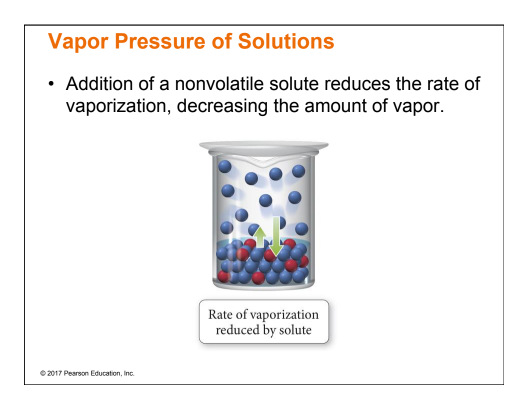
- 1. Write the given concentration as a ratio.
- 2. Separate the numerator and denominator.– Separate into the solute part and solution part.
- 3. Convert the solute part into the required unit.
- 4. Convert the solution part into the required unit.
- 5. Use the definitions to calculate the new concentration units.

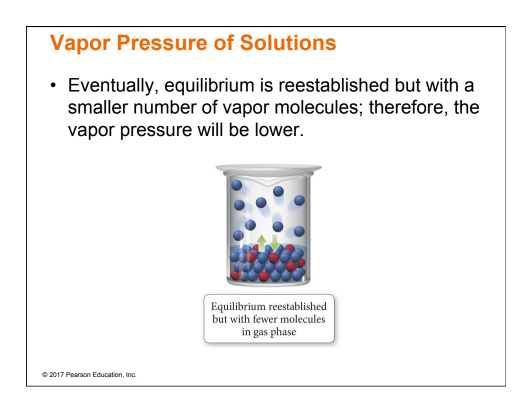
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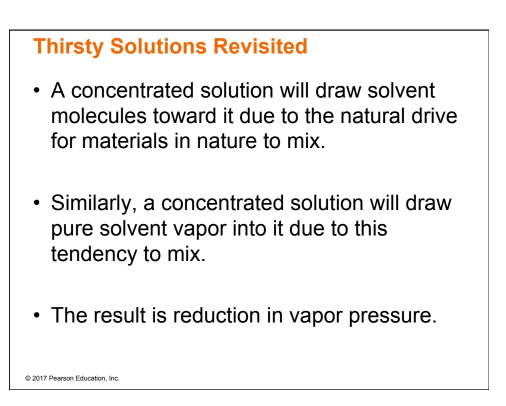
Colligative Properties

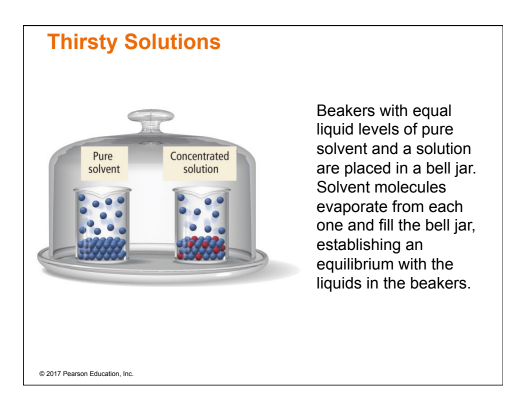
- **Colligative properties** are properties whose values depend only on the number of solute particles and not on what they are.
 - Value of the property depends on the concentration of the solution.
- The difference in the value of the property between the solution and the pure substance is generally related to the different attractive forces and solute particles occupying solvent molecules' positions.

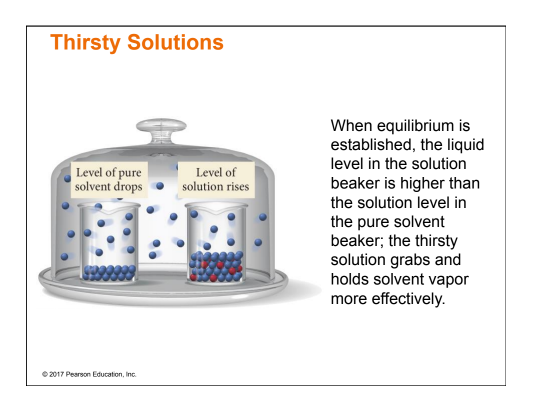


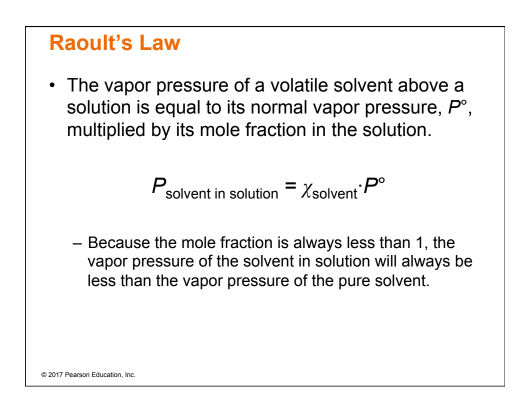


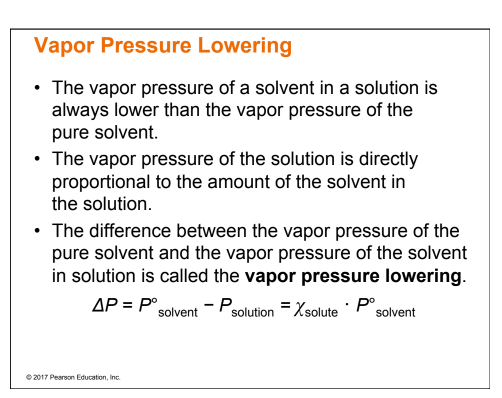












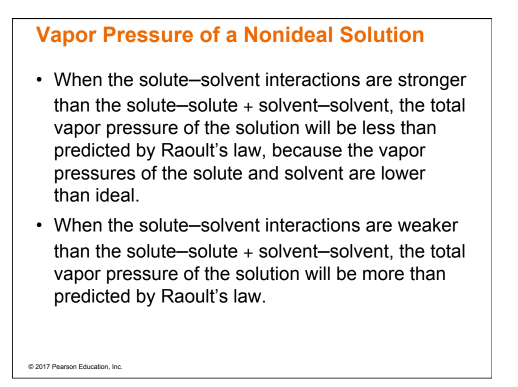
Raoult's Law for Volatile Solute • When both the solvent and the solute can evaporate, both molecules will be found in the vapor phase. • The total vapor pressure above the solution will be the sum of the vapor pressures of the solute and solvent. - For an ideal solution $P_{total} = P_{solute} + P_{solvent}$ • The solvent decreases the solute vapor pressure in the same way the solute decreased the solvent's. $P_{solute} = \chi_{solute} \cdot P^{\circ}_{solute}$ and $P_{solvent} = \chi_{solvent} \cdot P^{\circ}_{solvent}$

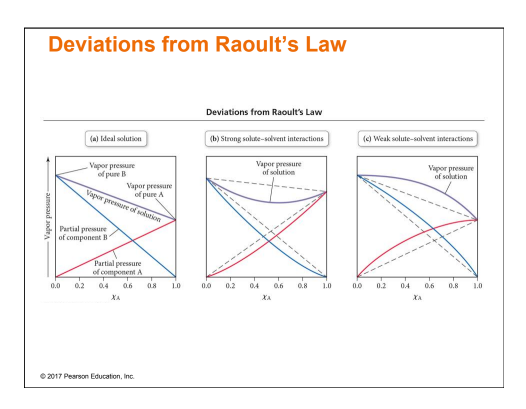
Ideal versus Nonideal Solutions

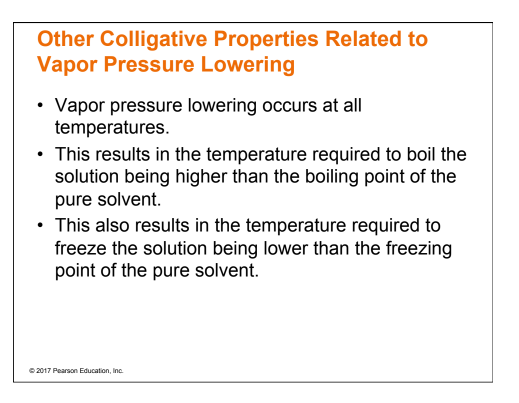
• In ideal solutions, the made solute–solvent interactions are equal to the sum of the broken solute–solute and solvent–solvent interactions.

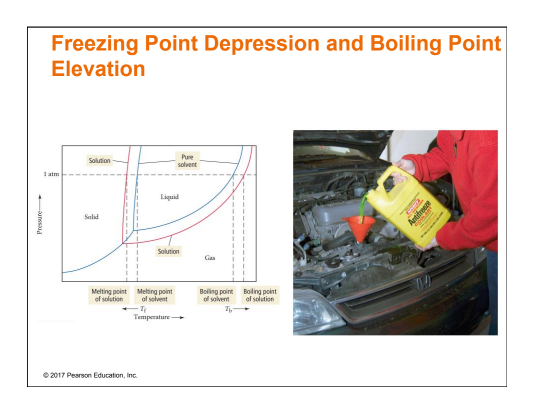
- Ideal solutions follow Raoult's law.

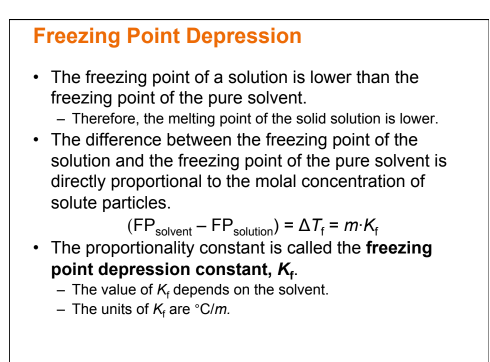
- Effectively, the solute is diluting the solvent.
- If the solute–solvent interactions are stronger or weaker than the broken interactions, the solution is nonideal.





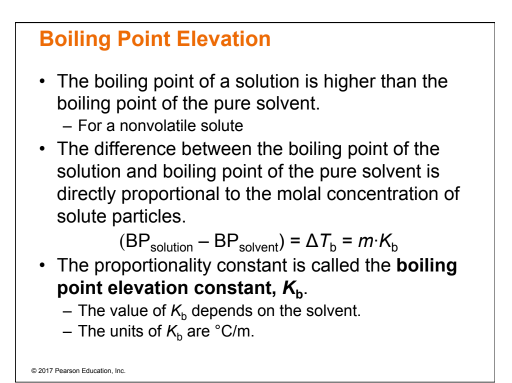






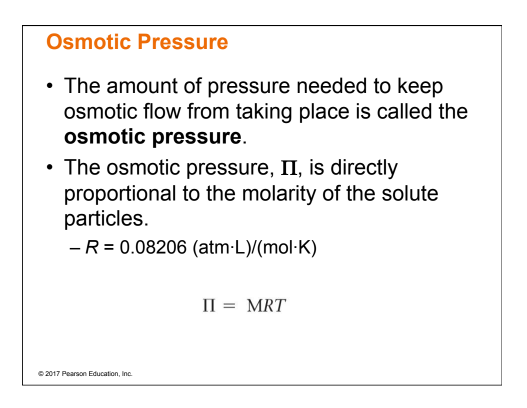
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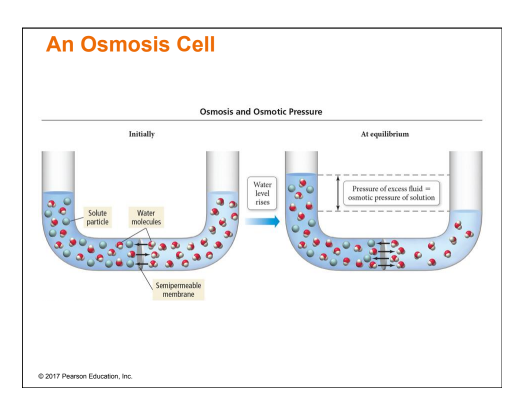
TABLE 13.8 Freezing Po	oint Depression and Boiling Po	int Elevation Co	nstants for Several Liquid Solver	nts
Solvent	Normal Freezing Point (°C)	K₁(°C/m)	Normal Boiling Point (°C)	К _b (°С/ <i>m</i>)
Benzene (C ₆ H ₆)	5.5	5.12	80.1	2.53
Carbon tetrachloride (CCl ₄)	-22.9	29.9	76.7	5.03
Chloroform (CHCl ₃)	-63.5	4.70	61.2	3.63
Ethanol (C ₂ H ₅ OH)	-114.1	1.99	78.3	1.22
Diethyl ether (C ₄ H ₁₀ O)	-116.3	1.79	34.6	2.02
Water (H ₂ O)	0.00	1.86	100.0	0.512

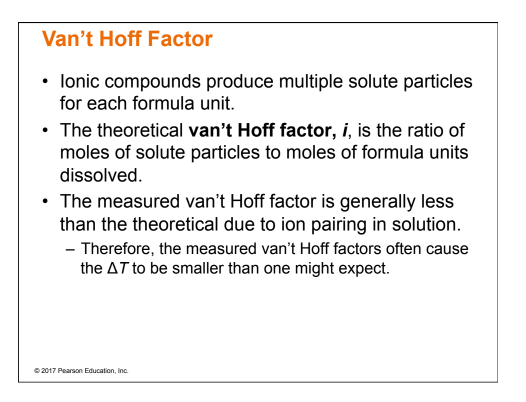


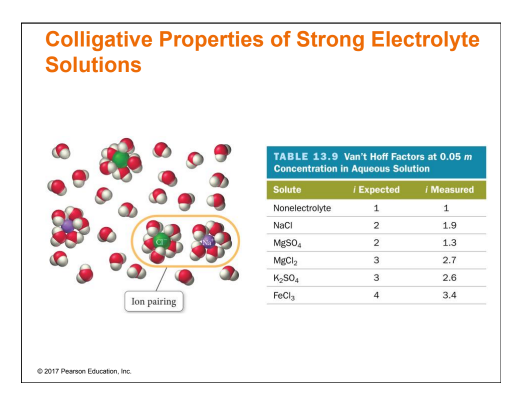
Osmosis

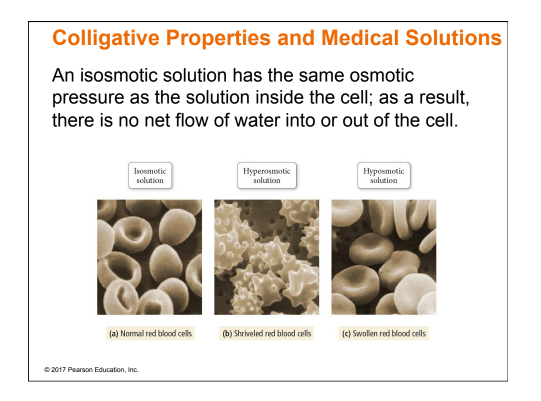
- **Osmosis** is the flow of solvent from a solution of low concentration into a solution of high concentration.
- The solutions may be separated by a semipermeable membrane.
- A semipermeable membrane allows solvent, but not solute, to flow through it.

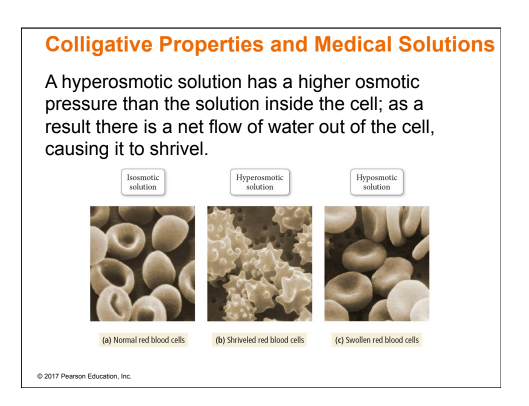


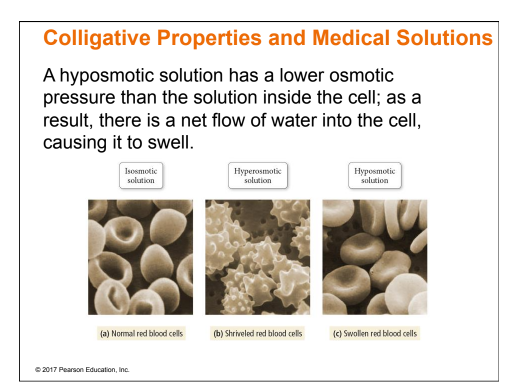












Mixtures
 Solutions = homogeneous Suspensions = heterogeneous, separate on standing Colloids = heterogeneous, do not separate on standing Particles can coagulate. Cannot pass through semipermeable membrane Hydrophilic Stabilized by attraction for solvent (water)
 Hydrophobic Stabilized by charged surface repulsions
 Show the Tyndall effect and Brownian motion.
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Types of	TABLE 13.10 Types of Colloidal Dispersions				
	Classification	Dispersing Substance (Solute-like)	Dispersing Medium (Solvent-like)		Example
	Aerosol	Liquid	Gas	N.	Fog (water droplets in air)
	Solid aerosol	Solid	Gas	1	Smoke (ash in air)
	Foam	Gas	Liquid	فلللن	Whipped cream (air bubbles in butterfat)
	Emulsion	Liquid	Liquid	ý	Milk (milk fat globules in water)
	Solid emulsion	Liquid	Solid		Opal (water in silica glass)
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