# Phases, changes of state

# Vapor pressure, boiling, and phase maps

A given substance will exist in the form of a solid, liquid, or gas, depending on the temperature and pressure. In this unit, we will learn what common factors govern the preferred state of matter under a particular set of conditions, and we will examine the way in which one phase gives way to another when these conditions change.



Water's changes of state contribute to the drama of nature. Seascape study with rain clouds - John Constable, 1825

# 1 Stability of a phase



Earlier in the morning, these droplets were tiny crystals of ice, but even though the air temperature is still around 0°C and will remain so all day, the sun's warmth has rendered them into liquid form, bound up by surface tension into reflective spheres. By late afternoon, most of the drops will be gone, their  $H_2O$  molecules now dispersed as a tenuous atmospheric gas.

[image: Crawford Wilson III]

Solid, liquid, and gas — these are the basic three states, or *phases*, in which the majority of small-molecule substances can exist. At most combinations of pressure and

temperature, only one of these phases will be favored; this is the phase that is most *thermodynamically stable* under these conditions.

### Origins of stability: why is one phase usually favored?

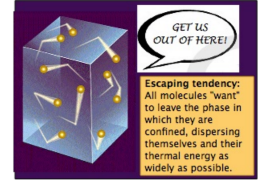
A proper explanation of why most substances have definite melting and boiling points needs to invoke some principles of thermodynamics and quantum mechanics. <u>See here</u> for a reasonably understandable explanation. Farther down on the same page is a more complete definition of "escaping tendency". A full explanation of this would go beyond the scope of what most students who see this lesson are familiar with, but the following greatly over-simplified explanation should convince you that it is something more than black magic:

All atoms and molecules at temperatures above absolute zero possess thermal energy that keeps them in constant states of motion. A fundamental law of nature mandates that this energy tends to spread out and be shared as widely as possible. Within a single molecular unit, this spreading and sharing can occur by dispersing the energy into the many allowed states of motion (translation, vibration, rotation) of the molecules of the substance itself. There are a huge number of such states, and they are quantized, meaning that they all require different amounts of thermal energy to come into action. Temperature is a measure of the intensity of thermal energy, so the higher the temperature, the greater will be the number of states that can be active, and the more extensively will the energy be dispersed among these allowed states.

In solids, the molecular units are bound into fixed locations, so the kinds of motion (and thus the number of states) that can be thermally activated is relatively small. Because the molecules of solids possess the lowest potential energies, solids are the most stable states at low temperatures. At the other extreme are gas molecules which are not only free to vibrate and rotate, but are in constant translational motion. The corresponding number of quantum states is hugely greater for gases, providing a nearlyendless opportunity to spread energy. But this can only happen if the temperature is high enough to populate this new multitude of states. Once it does, the gaseous state wins out by a landslide.

### Escaping tendency of a substance from a phase

Escaping tendency is more formally known as *free energy*, a fundamental concept that we explain in <u>this section</u> of the lesson set <u>Thermodynamics of</u> <u>Equilibrium</u>. Bear in mind also that changes in state always involve changes in enhalpy and internal energy; see <u>this section</u> of the energetics lesson for a quick overview. In much the same way that tea spreads out from a tea bag into the larger space of the water in which it is immersed, molecules that are confined within a phase (liquid, solid, or gas) will tend to spread themselves (and the thermal energy they carry with them) as widely as possible. This fundamental law of nature is manifested in what we will call the *escaping tendency* of the molecules from the phase. The escaping tendency is a quantity of fundamental importance in understanding all chemical equilibria and transformations. We need not define the term in a



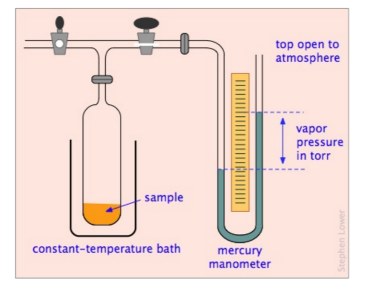
formal way at this point. What is important for now is how we can observe and compare escaping tendencies.

Think first of a **gas**: what property of the gas constitutes the best measure of its tendency to escape from a container? It does not require much reflection to conclude that the greater the *pressure* of the gas, the more frequently will its molecules collide with the walls of the container and possibly find their way through an opening to the outside. Thus the *pressure* confining a gas is a direct measure of the tendency of molecules to escape from a gaseous phase.

What about **liquids and solids**? Although we think of the molecules of condensed phases as permanently confined within them, these molecules still possess some thermal energy, and there is always a chance that one that is near the surface will occasionally fly loose and escape into the space outside the solid or liquid.

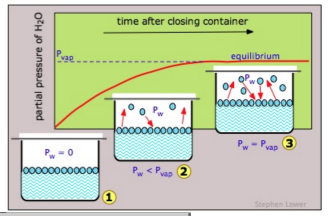
### Vapor pressure

We can observe the tendency of molecules to escape into the gas phase from a solid or liquid by placing the substance in a closed, evacuated container connected to a manometer for measuring gas pressure.



will rise as molecules escape from the substance and enter the vapor phase. But at the same time, some of the vapor molecules will "escape" back into the liquid phase (2). But because this latter process is less favorable (at the particular temperature represented here),  $P_W$  continues to rise. Eventually a balance is reached between the two processes (3), and  $P_W$  eventually stabilizes at a fixed value  $P_{vap}$  that depends on the substance and on the temperature and is known as the **equilibrium vapor** 

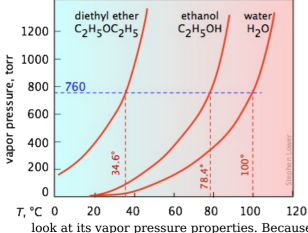
*pressure*, or simply as the "vapor pressure" of the liquid or solid. The vapor pressure is a direct measure of the escaping tendency of molecules from a condensed state of matter.



Note carefully that if the container is left open to the air, it is unlikely that many of the molecules in the vapor phase will return to the liquid phase. They will simply escape from the entire system and the partial pressure of water vapor  $P_W$  will never reach  $P_{vap}$ ; the liquid will simply *evaporate* without any kind of equilibrium ever being achieved.

#### Vapor pressure plots and boiling points

The escaping tendency of molecules from a phase always increases with the temperature; therefore the vapor pressure of a liquid or solid will be greater at higher temperatures. As the figure shows, the variation of the vapor pressure with the temperature is not linear.



look at its vapor pressure properties. Because the vapor pressure of water varies greatly over the range of temperatures in which the liquid can exist, the illustration shows two plots. The larger plot covers the lowest temperatures, while the inset shows the complete range of pressure values. Note particularly that

- The *normal boiling point* is the temperature at which the vapor pressure is the same as that of the standard atmosphere, 760 torr.
- The boiling point at any other pressure can be found by dropping a vertical line from the curve to the temperature axis.
- As seen on the inset plot, the vapor pressure curve of water ends at the *critical point*.

### **Relative humidity**

The vapor pressure of water at 22°C is about 20 torr, or around 0.026 atm (2.7 KPa). This is the partial pressure of  $H_2O$  that will be found in the vapor space

such as the three shown here. Take special note of how boiling points can be found from these plots. You will recall that the *normal boiling point* is the temperature at which the liquid is in equilibrium with its vapor at a partial pressure of 1 atm (760 torr). Thus the intercepts of each curve with the blue dashed 760-torr line indicate the normal boiling points of each liquid. Similarly, you can easily estimate the boiling points these liquids would have in Denver, Colorado where the atmospheric pressure is 630 torr by simply constructing a horizontal line corresponding to this pressure.

It's important that you be able to interpret vapor pressure plots

#### Vapor pressure of water

torr 250 10.000 atm atm MP critical point 374 K, 218 atm 200 20 9.000 KPa 8,000 150 10 1000 7,000 100 10 6,000 8 800 50 5.000 6 600 4,000 0 200 300 400 0 3,000 4 400 2.000 200 pressure 1,000 normal boiling point 0 0 50 100 150 200 temperature, K

The great importance of  $H_2O$  in our world merits a more detailed

within a closed container of water at this temperature; the air in this space is said to be *saturated* with water vapor.

Humid air is sometimes described as "heavy", but this is misleading; the average molar mass of dry air is 29, but that of water is only 18, so humid air is actually *less* dense. (Review Dalton's Law of partial pressures <u>here</u>!) The feeling of "heaviness" probably relates to the reduced ability of perspiration to evaporate in humid air.

In ordinary air, the partial pressure of water vapor is normally less than its saturation or equilibrium value. The ratio of the partial pressure of  $H_2O$  in the air to its (equilibrium) vapor pressure at any given temperature is known as the *relative* 

*humidity*. Water enters the atmosphere through evaporation from the ocean and other bodies of water, and from watersaturated soils. The resulting vapor tends to get dissipated and diluted by atmospheric circulation, so the relative humidity rarely reaches 100 percent. When it does and the weather is warm, we are very uncomfortable because vaporization of water from the skin is inhibited; if the air is already saturated with water, then there is no place for our perspiration to go, other than to drip down our face.

#### Why the indoor air is so dry in the winter

Because the vapor pressure increases with temperature, a parcel of air containing a fixed partial pressure of water vapor will have a larger relative humidity at low temperatures than at high temperatures. Thus when cold air enters a heated house, its water content remains unchanged but the relative humidity drops. In climates with cold winters, this promotes increased moisture loss from house plants and from mucous membranes, leading to wilting of the former and irritation of the latter.

#### Problem Example 1

The vapor pressure of water is 3.9 torr at  $-2^{\circ}\text{C}$  and 20 torr at  $22^{\circ}\text{C}$ . What will be the relative humidity inside a house maintained at  $22^{\circ}\text{C}$  when the outside air temperature is  $-2^{\circ}\text{C}$  and the relative humidity is 70%?

Solution:

At 70 percent relative humidity, the partial pressure of the  $-2^{\circ}$  air is  $(0.7 \times 3.9 \text{ torr}) = 2.7 \text{ torr}$ . When this air enters the house, its relative humidity will be (2.7 torr)/(20 torr) = 0.14 or 14%.



In the evening, especially on clear nights, solid objects (even spider webs!) lose heat to the sky more rapidly than does the air. It is often important to know what temperature such objects must drop to so that atmospheric moisture will condense out on them.

The *dew point* is the temperature at which the relative humidity is 100 percent — that is, the temperature at which the vapor pressure of water becomes equal to its partial pressure at a given [higher] temperature and relative humidity. [image]

For water to condense directly out of the atmosphere as rain, the air must be at or below the dew point, but this is not of itself the only requirement for the formation of rain, as we will see shortly.

### Vapor pressures of solid hydrates

Many solid salts incorporate water molecules into their crystal lattices; the resulting compounds are known as *hydrates*. These solid hydrates possess definite vapor pressures which correspond to equilibrium between the hydrated and anhydrous compounds and water vapor. For example:

 $SrCl_2 \cdot 6H_2O(s) \rightarrow SrCl_2(s) + 6H_2O(g)$ 

The vapor pressure of this hydrate is 8.4 torr at 25°C. Only at this unique partial pressure of water vapor can the two solids coexist at 25°C. If the partial pressure of water in the air is greater than 8.4 torr, a sample of anhydrous  $SrCl_2$  will absorb moisture from the air and change into the hydrate.

#### **Problem Example 2**

What will be the relative humidity of air in an enclosed vessel containing solid SrCl<sub>2</sub>·6H<sub>2</sub>O at 25°C? Solution:

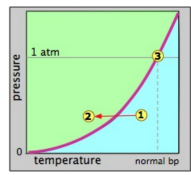
What fraction of the vapor pressure of water at this temperature (23.8 torr) is the vapor pressure of the hydrate (8.4 torr)? Expressed in percent, this is the relative humidity.

If the partial pressure of  $H_2O$  in the air is less than the vapor pressure of the hydrate, the latter will tend to lose moisture and revert back to its anhydrous form. This process is sometimes accompanied by a breakup of the crystal into a powdery form, an effect known as *efflorescence*.

# 2 Condensation and boiling: nucleation

Evaporation and boiling of a liquid, and condensation of a gas (vapor) are such ordinary parts of our daily life that we hardly give them a thought. Every time we boil water to make a pot of tea and see the cloud of steam above the teapot, we are observing this most common of all phase changes. How can we understand these changes in terms of vapor pressure?

This plot of vapor-pressure as a function of temperature can represent water or any other liquid. When we say this is a vapor-pressure plot, we mean that each point on the curve represents a combination of temperature and vapor pressure at which the liquid (green) and the vapor (blue) can coexist. Thus at the normal boiling point, defined as the temperature at which the vapor pressure is 1 atm, the state of the system corresponds to the point labeled **3**.



Suppose that we select an arbitrary point 1 at a temperature and pressure at which only the gaseous state is stable. We then decrease the temperature so as to move the state point toward point 2 in the liquid region. When the state point falls on the vapor pressure line, the two phases can coexist and we would expect some liquid to condense. Once the state point moves to the left of the vapor

pressure line, the substance will be entirely in the liquid phase. This is supposedly what happens when "steam" (actually tiny water drops) forms above a pot of boiling water.

The reverse process should work the same way: starting with a temperature in the liquid region, nothing happens until we reach the vapor pressure line, at which point the liquid begins to change into vapor. At higher temperatures, only the vapor remains.

This is the *theory*, but it is not complete: the *fact* is that a vapor will generally *not* condense to a liquid at the boiling point (also called the condensation point or dew point), and a liquid will generally *not* boil at its boiling point.

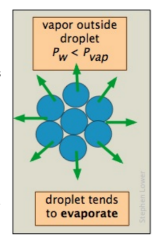
### **Bubbles and drops**

The reason for the discrepancy is that the vapor pressure, as we normally use the term and as it is depicted by the liquidvapor line on the phase diagram, refers to the partial pressure of vapor in equilibrium with a liquid whose surface is reasonably flat, as it would be in a partially filled container. In a drop of liquid or in a bubble of vapor within a liquid, the surface of the liquid is not flat, but curved. For drops or bubbles that are of reasonable size, this does not make much difference, but these drops and bubbles must grow from smaller ones, and these from tinier ones still. Eventually, one gets down to the primordial drops and bubbles having only a few molecular dimensions, and it is here that we run into a problem: this is the problem of **nucleation** — the formation and growth of the first tiny drop (in the vapor) or of a bubble (in a liquid).

### The nucleation problem

The vapor pressure of a liquid is determined by the attractive forces that act over a 180° solid angle at the surface of a liquid. In a very small drop, the liquid surface is curved in such a way that each molecule experiences fewer nearest-neighbor attractions than is the case for the bulk liquid. The outermost molecules of the liquid are bound to the droplet less tightly, and the drop has a larger vapor pressure than does the bulk liquid. If the vapor pressure of the drop is greater than the partial pressure of vapor in the gas phase, the drop will evaporate. Thus it is highly unlikely that a droplet will ever form within a vapor as it is cooled.

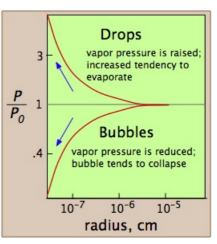
> A bubble, like a drop, must start small and grow larger, but there is a difficulty here that is similar to the one with bubbles. A bubble is a hole in a liquid; molecules at the liquid boundary are curved inward, so that they experience nearest-neighbor attractions over a solid angle greater than 180°. As a consequence, the vapor pressure of the liquid facing into a bubble  $P_{W}$  is always less than that of the bulk



liquid  $P_{w}$  at the same temperature. When the bulk liquid is at its boiling point (that is, when its vapor pressure is 1 atm), the pressure of the vapor within the bubble will be less than 1 atm, so the bubble will tend to collapse. Also, since the bubble is formed within the liquid, the hydrostatic pressure of the overlaying liquid will add to this effect. For both of these reasons, a liquid will not boil until the temperature is raised slightly above the boiling point, a phenomenon known as *superheating*. Once the boiling begins, it will continue to do so at the liquid's proper boiling point.

These plots show how, in the case of water, the vapor pressure of a very small bubble or drop

varies with its radius of curvature; the quantity being plotted is the ratio of the actual vapor pressure *P* to  $P_{o'}$  the vapor pressure of a flat surface.



### **Condensation of a liquid**

vapor above liquid

 $P_w = P_{vap}$ 

bubble tends to

collapse

vapor inside bubble

Pw < Pvap

If the tiniest of drops are destined to self-destruct, why do vapors ever condense, and why does it rain?

- If you cool a vapor in a container, condensation takes place not within the vapor itself, but on the inner surface of the container. What happens here is that intermolecular attractions between the solid surface will cause vapor molecules to adsorb to the surface and stabilize the incipient drop until it grows to a size at which it can be self-sustaining. This is the origin of the condensation on the outside of a cool drink, or of the dew that appears on the grass.
- In the case of the cloud of steam you see over the boiling water, the first few droplets form on tiny dust particles in the air the ones you can see by scattered light when a sunbeam shines through a darkened room.

This Wikipedia article describes how cloud condensation nuclei form. The conditions under which cloud droplets coalesce into rain is not thoroughly understood. For a readable overview, see here.

**Clouds and precipitation** In the region of the atmosphere where rain forms there are large numbers of solid particles, mostly of microscopic size. Some of these are particles of salt produced by evaporation of spray from the ocean surface. Many condensation nuclei are of biological origin; these include bacteria, spores, and particles of ammonium sulfate. There is volcanic and meteor dust, and of course there is dust and smoke due to the activities of humans. These particles tend to adsorb water vapor, and some may even dissolve to form a droplet of concentrated solution. In either case, the vapor



pressure of the water is reduced below its equilibrium value, thus stabilizing the aggregate until

it can grow to self-sustaining size and become fog, rain, or snow.

This, by the way, is why fog is an irritant to the nose and throat; each fog droplet carries within it a particle of dust or (in air polluted by the burning of sulfur-containing fossil fuels) a droplet of sulfuric acid, which it efficiently deposits on your sensitive mucous membranes. If you own a car which is left outside on a foggy night, you may have noticed how dirty the windshield is in the morning.

### Superheating and boiling of liquids

What is the difference between the evaporation and boiling of a liquid? When a liquid evaporates at a temperature below its boiling point, the molecules that enter the vapor phase do so directly from the surface. When a liquid boils, bubbles of vapor form in the interior of the liquid, and are propelled to the surface by their lower density (buoyancy). As they rise, the diminishing hydrostatic pressure causes the bubbles to expand, reducing their density (and increasing their buoyancy) even more.

But as we explained above, getting that first bubble to form and survive is often sufficiently difficult that liquids commonly superheat before the begin to boil.

In large quantities, superheated liquids can be very dangerous, because the introduction of an impurity (such as release of

If you have had experience in	
an organic chemistry	
laboratory, you probably	
know this as "bumping", and	
have been taught to take	
precautions against it.	

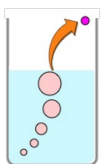
an air bubble from the container surface) or even a mechanical disturbance can trigger nucleation and cause boiling to occur suddenly and almost explosively.

#### Exploding water in the microwave

Many people have been seriously burned after attempting to boil water in a microwave oven, or after having added powdered material such as instant coffee to such water. When water is heated on a stove, the bottom of the container superheats only the thin layer of water immediately in contact with it, producing localized "microexplosions" that you can hear just before regular smooth boiling begins; these bubbles quickly disperse and serve as nucleation centers for regular boiling.

In a microwave oven, however, the energy is absorbed by the water itself, so that the entire bulk of the water can become superheated. If this happens, the slightest disturbance can produce an explosive flash into vapor. <u>See here</u> for more and for some suggestions on how to avoid this danger.

#### Sublimation

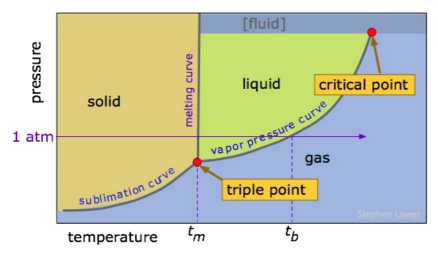


Some solids have such high vapor pressures that heating leads to a substantial amount of direct vaporization even before the melting point is reached. This is the case for solid iodine, for example.  $I_2$  melts at 115°C and boils at 183°C, is easily

sublimed at temperatures around 100°C. Even ice has a measurable vapor pressure near its freezing point, as evidenced by the tendency of snow to evaporate in cold dry weather.

There are other solids whose vapor pressure overtakes that of the liquid before melting can occur. Such substances sublime without melting; a common example is solid carbon dioxide ("Dry Ice") at 1 atm (see the  $CO_2$  phase diagram below).

# 3 Phase maps



The temperatures and pressures at which a given phase of a substance is stable (that is, from which the molecules have the lowest escaping tendency) is an important property of any substance. Because both the temperature and pressure are factors, it is customary to plot the regions of stability of the various phases in P-T coordinates, as in this generic **phase diagram** (or phase map) for a hypothetical substance.

Because pressures and temperatures can vary over very wide ranges, it is common practice to draw phase diagrams with non-linear or

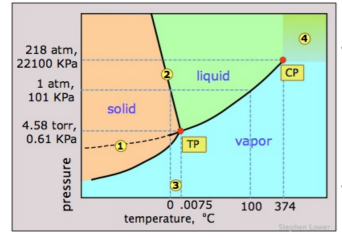
distorted coordinates. This enables us to express a lot of information in a compact way and to visualize changes that could not be represented on a linearly-scaled plot.

It is important that you be able to interpret a phase map, or alternatively, construct a rough one when given the appropriate data. Take special note of the following points:

- 1. The three colored regions on the diagram are the ranges of pressure and temperature at which the corresponding phase is the only stable one.
- 2. The three lines that bound these regions define all values of (*PT*) at which *two* phases can coexist (i.e., be in equilibrium). Notice that one of these lines is the *vapor pressure curve* of the *liquid* as described above. The "*sublimation curve*" is just a vapor pressure curve of the *solid*. The slope of the line depends on the difference in density of the two phases.
- 3. In order to depict the important features of a phase diagram over the very wide range of pressures and temperatures they encompass, the axes are not usually drawn to scale, and are usually highly distorted. This is the reason that the "melting curve" looks like a straight line in most of these diagrams.
- 4. Where the three named curves intersect, all three phases can coexist. This condition can only occur at a unique value of (P,T) known as the *triple point*. Since all three phases are in equilibrium at the triple point, their vapor pressures will be identical at this temperature.
- 5. The line that separates the liquid and vapor regions ends at the *critical point*. At temperatures and pressures greater than the critical temperature and pressure, no separate liquid phase exists. We refer to this state simply as a *fluid*, although the term *supercritical liquid* is also commonly used.

The best way of making sure you understand a phase diagram is to imagine that you are starting at a certain temperature and pressure, and then change just one of these parameters, keeping the other constant. You will be traversing a horizontal or vertical path on the phase map, and there will be a change in state every time your path crosses a line. Of special importance is the horizontal path (shown by the blue line on the diagram above) corresponding to a pressure of 1 atmosphere; this line defines the *normal melting and boiling temperatures* of a substance.

### **Phase map of Water**

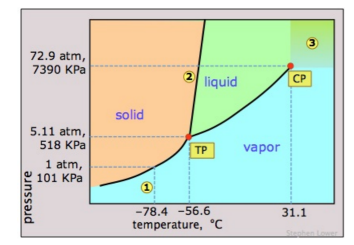


Notice the following features of this very important phase diagram:

- The slope of the line **2** separating the solid and liquid regions is negative; this reflects the unusual property that the density of the liquid is greater than that of the solid, and it means that the melting point of ice decreases as the pressure increases. Thus if ice at 0°C is subjected to a high pressure, it will find itself above its melting point and it will melt. (Contrary to what is sometimes said, however, this is not the reason that ice melts under the pressure of ice skates or skis, providing a lubricating film which makes these modes of transportation so enjoyable. The melting in these cases arises from frictional heating.)
- The dashed line **1** is the extension of the liquid vapor pressure line below the freezing point. This represents the vapor pressure of *supercooled* water
  - a state of water which can temporarily exist down to about -20°C. (If you live in a region subject to "freezing rain", you

will have encountered supercooled water!)

- 3 The triple point (TP) of water is just  $0.0075^{\circ}$  above the freezing point; only at this temperature and pressure can all three phases of water coexist indefinitely.
- 4 Above the critical point (CP) temperature of 374°C, no separate liquid phase of water exists.

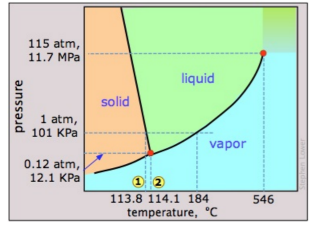


### Phase map of Carbon dioxide

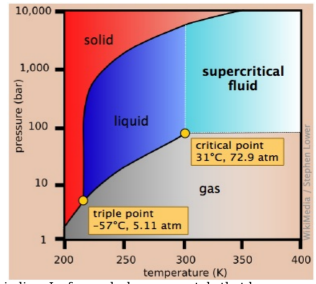
Solid carbon dioxide, commonly known as Dry Ice, is widely used as a refrigerant. The phase diagram shows why it is "dry". The triple point pressure is at 5.11 atm, so below this pressure, liquid  $CO_2$  cannot exist; the solid can only sublime directly to vapor. Gaseous carbon dioxide at a partial pressure of 1 atm is in equilibrium with the solid at 195K (-79 °C, **1**); this is the *normal sublimation temperature* of carbon dioxide. The surface temperature of dry ice will be slightly less than this, since the partial pressure of  $CO_2$  in contact with the solid will usually be less than 1 atm. Notice also that the critical temperature of  $CO_2$  is only 31°C. This means that on a very warm day, the  $CO_2$  in a fire extinguisher will be entirely vaporized; the vessel must therefore be strong enough to withstand a pressure of 73 atm.

This view of the carbon dioxide phase map employs a logarithmic pressure scale and thus encompasses a much wider range of pressures, revealing the upper boundary of the fluid phase (liquid and supercritical). **Supercritical carbon dioxide** (CO<sub>2</sub> above its critical temperature) possesses the solvent properties of a liquid and the penetrating properties of a gas; one major use is to remove caffeine from coffee beans.

### Phase diagram of iodine



temperatures of its triple point 1 and melting point 2.



Elemental iodine,  $I_2$ , forms dark gray crystals that have an almost metallic appearance. It is often used in chemistry classes as an example of a solid that is easily sublimed; if you have seen such a demonstration or experimented with it in the lab, its phase diagram might be of interest.

The most notable feature of iodine's phase behavior is the very small difference (less than a degree) between the

Contrary to the impression many people have, there is nothing really special about iodine's tendency to sublime, which is shared by many molecular crystals including ice and naphthalene ("moth crystals".) The vapor pressure of iodine at room temperature is really quite small — only about 0.3 torr (40 Pa). The fact that solid iodine has a strong odor and is surrounded by a purple vapor in a closed container is mainly a consequence of its strong ability to absorb green light (this leaves blue and red which make purple) and the high sensitivity of our noses to its vapor. [image]

### Phase diagram of sulfur

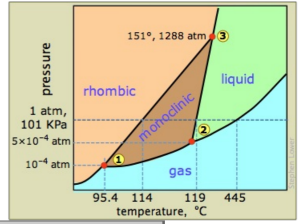
Sulfur exhibits a very complicated phase behavior that has puzzled chemists for over a century; what you see here is the greatly simplified phase map shown in most textbooks. The difficulty arises from the tendency of  $S_8$  molecules to break up into chains (especially in the liquid above 159°C) or to rearrange into rings of various sizes ( $S_6$  to  $S_{20}$ ). Even the vapor can contain a mixture of species  $S_2$  through  $S_{10}$ .



The phase diagram of sulfur contains a new feature: there are two solid phases, *rhombic* and *monoclinic*. The names refer to the crystal structures in which the  $S_8$  molecules arrange

themselves. This gives rise to three triple points, indicated by the numbers on the diagram. [Question: which three phases can *never* coexist?]

When rhombic sulfur (the stable low-temperature phase) is heated slowly, it changes to the monoclinic form at 114°C, which then melts at 119°. But if the monoclinic form is heated rapidly the molecules do not have time to rearrange themselves, so the rhombic arrangement persists as a metastable phase until it melts at 119-120°.



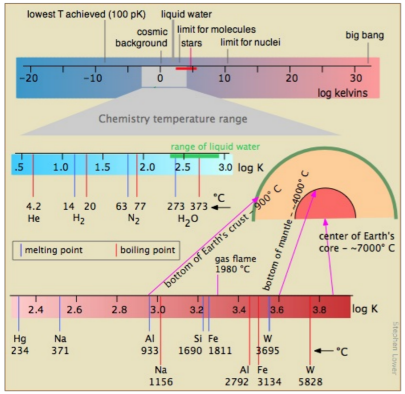
Formation of more than one solid phase is not uncommon — in fact, if one explores into the very high pressures (see below), it seems to be the rule.

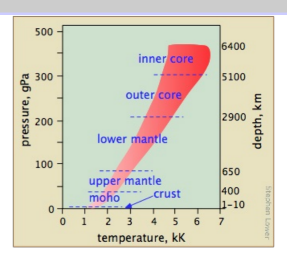
# 4 Phases at the extremes

We tend to think of the properties of substances as they exist under the conditions we encounter in everyday life, forgetting that most of the matter that makes up our world is situated inside the Earth, where pressures and temperatures are orders of magnitude higher. Geochemists and planetary scientists need to know about the phase behavior of substances at high temperatures and pressures in order to develop useful models to test their theories about the structure and evolution of the Earth and of the solar system.

What ranges of temperatures and pressures are likely to be of interest — and more importantly, are experimentally accessible?

# Extreme temperatures





This diagram shows several scales (all of which, please note, are logarithmic) that cover respectively

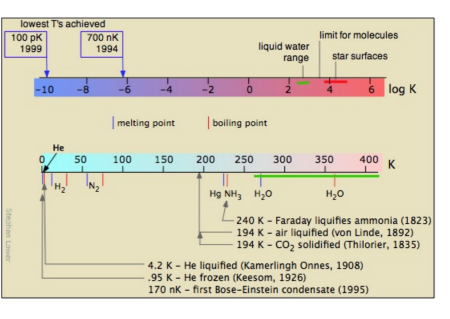
- the temperature range for the universe;
- the low temperatures of importance to chemistry (note the green line indicating the temperatures at which liquid water can exist);
- the higher temperatures, showing the melting and boiling points of several elements for reference.

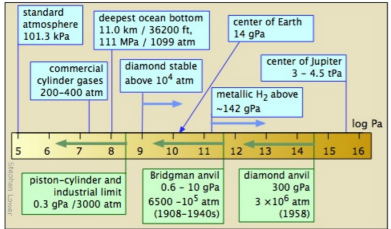
The highest temperatures that can be produced in the laboratory are achieved (but only for very short time intervals) by light pulses from laser or synchrotron radiation. The study of **low temperatures** is limited by the laws of physics that prohibit reaching absolute zero. But the fact that there is no limit to how close one can approach 0 K has encouraged a great deal of creative experimentation.

Some references:

- detailed  $\underline{\operatorname{timeline}}$  of low-temperature technology
- <u>Carl von Linde</u> (liquifaction of air)
- <u>Heike Kammerlingh Onnes</u> (1913 Nobel <u>Prize in physics</u>)
- Bose-Einstein condensation

# Phase studies at high pressures.

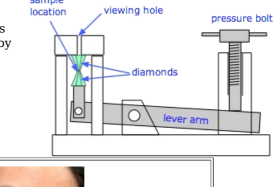


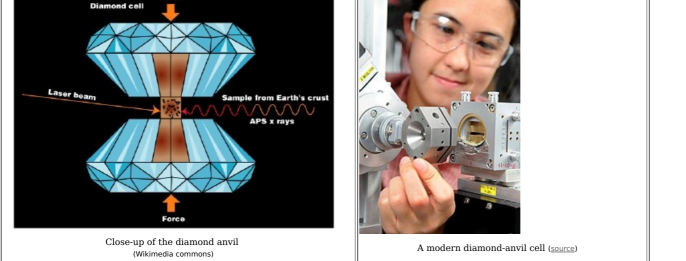


The study of matter at high pressures is not an easy task. The general techniques were pioneered between 1908-1960 by <u>P.W.</u> <u>Bridgman</u> of Harvard University, whose work won him the 1946 Nobel Prize in physics. The more recent development of the diamond anvil cell has greatly extended the range of pressures attainable and the kinds of observations that can be made. Shock-wave techniques have made possible the production of short-lived pressures in the tPa range.

#### **Diamond anvil cell**

The *diamond anvil cell*, developed in 1959, produces the highest static pressures available in the laboratory. Diamond is used because of its extreme hardness and its transparency, which allows the sample to be viewed through a microscope. Thrust generated by the pressure bolt is transmitted to the diamonds through a lever arm, piston, and rocker mounts. The cell itself is a hole in a thin sheet of metal placed between the diamond surfaces; the metal serves as a gasket to prevent the contents of the cell from being squeezed out. (Illustration based on one in *Scientific American*, April 1984, pp 54-62.)





These studies have revealed that many molecular substances such as hydrogen and water change to solid phases having melting points well above room temperature at very high pressures; there is a solid form of ice that remains frozen even at 100°C. At still higher pressures, many of these substances become metals. It is believed that much of the inner portion of the largest planets consists of metallic hydrogen — and, in fact, that all substances can become metallic at sufficiently high pressures.

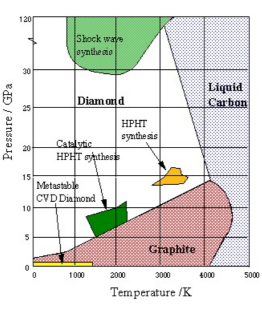
### Phase diagram of carbon: diamond and graphite

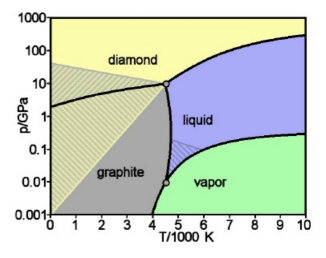
**Graphite** is the stable form of solid carbon at low pressures; diamond is only stable above about  $10^4$  atm. But once it is in this form, the rate at which diamond converts back to graphite is immeasurably slow under ordinary environmental conditions; there is simply not enough thermal energy available to break all of those carbon-carbon bonds. So the diamonds we admire in jewelry and pay dearly for are said to be . [image: WikiMedia]

### Synthetic diamonds

In order to transform graphite into diamond at a reasonable rate, a pressure of 200,000 atm and a temperature of about 4000 K would be required.

Since no apparatus can survive these conditions, the process, known as *high pressure high temperature synthesis* (HPHT) is carried out commercially at 70,000 atm and 2300 K in a solution of molten nickel, which also acts as a catalyst. Traces of Ni in the finished product

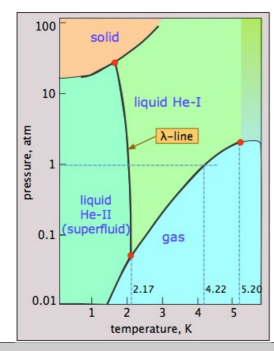




serve to distinguish synthetic diamonds from natural ones. But most synthetic diamonds are too small (only a few millimeters) and too flawed for gem quality, and are used mainly to fabricate industrial grinding and cutting tools.

More recently, thin diamond films have become important for engineering applications and semiconductor fabrication. These are most commonly made by condensation of gaseous carbon onto a suitable substrate (chemical vapor deposition, CVD). The conditions under which synthetic diamonds are made are depicted on the above phase diagram from Bristol University [link].

### Helium phase weirdness



Helium is unique in that quantum phenomena, which normally apply only to tiny objects such as atoms and electrons, extend to and dominate its macroscopic properties. A glance at the phase diagram of  $\mathrm{He}^4$  reveals some of this quantum weirdness. The main points to notice are

- helium can be frozen only at high pressures;
- the solid and gas cannot coexist (be in equilibrium) under any conditions;
- there are *two liquid phases*, helium I (an ordinary liquid) and helium II (a highly unusual *ordered liquid*); the  $\lambda$  (*lambda*) line represents the (*P*,*T*) values at which the two phases can coexist.
- helium-II behaves as a *superfluid* essentially a *quantum liquid*.

**Why can't liquid helium freeze at low pressures?** The low mass of the He atoms and their close confinement in the solid provides them with a very high *zero-point energy* (the Heisenberg uncertainty principle in action!) that allows them to vibrate with such amplitude that they overcome the dispersion forces that would otherwise hold the solid together, thus keeping the atoms too separated to form a solid. Only by applying a high pressure (25 atm) can this effect be overcome.

#### Helium-II, quantum liquids and superfluidity

We usually need quantum theory only to describe the properties of tiny objects such as electrons, but with liquid He-II, quantum effects extend to the macroscopic scale of the bulk liquid. He<sup>4</sup> atoms (99.99+ percent of natural helium) are *bosons*, which means that at low temperatures, they can all occupy the same quantum state. (All other normal atoms are *fermions* and are subject to the Pauli exclusion principle.)

#### All together, now

Objects that occupy the same quantum state all possess the same momentum. Thus when one atom moves, they all move together. In a sense, this means that the entire bulk of liquid He-II acts as a single entity. This property, known as *superfluidity*, gives rise to a number of remarkable effects, most notably:

- The liquid can flow through a narrow channel without friction, up to a critical velocity that depends on the ratio of flow rate to channel width;
- When placed in a container, they form a film that climbs up the walls and down the outside, seeking the same level outside the container as inside if this is possible;
- A small molecule dissolved in helium-II behaves the same as it would in a vacuum. [PhysicsWorld article].

In liquid helium-II, only about 10% of the atoms are in such a state, but it is enough to give the liquid some of the weird properties of a quantum liquid.

He<sup>3</sup> exhibits similar properties, but owing to its low natural abundance, it was not extensively studied until the 1940s when large amounts became available as a byproduct of nuclear weapons manufacture. Finally, in 1996, its superfluidity was observed at a temperature of 2 nK. Although He<sup>3</sup> atoms are fermions, only those that pair up (and thus assume the properties of bosons) give rise to superfluidity.

For more on superfluidity, see this <u>U. of Alaska site</u>, <u>this Wikipedia article</u>, this <u>Scientific American site</u> (with video) Another interesting video: <u>here</u>

### Water at high pressures

Water, like most other substances, exhibits many solid forms at higher pressures. So far, fifteen distinct ice phases have been identified; these are designated by Roman numerals ice-I through ice-XV. Ice-I can exist in two modifications; the crystal lattice of ice-Ic is cubic, while that of <u>ice-Ih</u> is hexagonal. The latter corresponds to the ordinary ice that we all know. It's interesting to note that several high-pressure phases of ice can exist at temperatures in excess of 100°C.

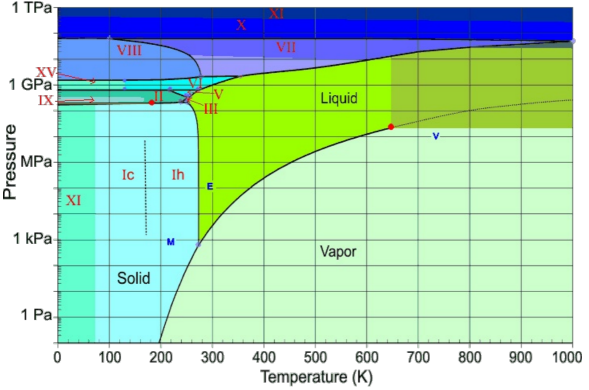


Image from Martin Chaplin's LSBU site

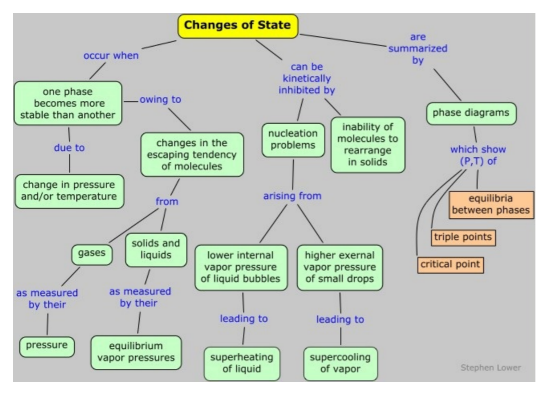
<u>Martin Chaplin's LSBU site</u> explains this diagram in great detail, and has links to separate pages that describe each of the ice phases. Another interesting page on this site deals with <u>supercooled "glassy" states</u> of water.

# What you should be able to do

Make sure you thoroughly understand the following essential ideas which have been presented above. It is especially imortant that you know the precise meanings of all the green-highlighted terms in the context of this topic.

- Describe what is meant by the *escaping tendency* of molecules from a solid, liquid, or gas. What experimentally-observable quantity serves as its measure?
- The terms "*vapor pressure*" and "*pressure of the vapor* above a solid or liquid" are easily confused. Explain the difference between them, and state under what conditions they will have identical values.
- Define *relative humidity* and calculate its value, given the partial pressure of water vapor and a suitable vapor pressure table or plot for water.
- Explain the difference between evaporation and boiling, and why liquids may not begin to boil until the temperature exceeds the boiling point.
- Given a *phase diagram* of a pure substance, label all of the lines and the regions they enclose, identify the *normal melting and boiling points*, the *triple point* and the *critical point*, and state the physical significance of the latter two.
- Conversely, sketch out a properly-labeled phase diagram for a pure substance, given the parameters mentioned above, along with information about the relative densities of the solid and liquid phases.

# **Concept map**



 $\ensuremath{\mathbb{C}}$  2009-2017 by Stephen Lower - last modified 2017-10-23

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