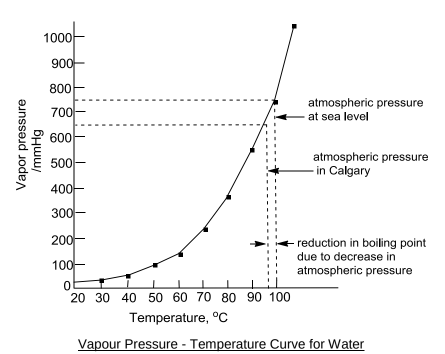
BOILING POINT DETERMINATION

The physical properties of a compound, such as melting point and boiling point can provide useful information which can help in the identification of a sample or to establish its purity. Since the boiling point of an unknown sample under the same conditions (*e.g.* same pressure) is a constant, a measured boiling point can be compared to known values (e.g. literature value or the measured value of a known sample). Since this used to be a very important method, there are tables and collections of boiling point data available and the boiling point at sea level (as a standard) is one of the physical properties of a compound that is commonly recorded and reported (*e.g.* see Wikipedia entry for ethanol)

Background

If a sample of a liquid is placed in an otherwise empty space, some of it will vapourise, and the pressure in the space above the liquid will rise to some constant value. The pressure under these conditions is due entirely to the vapour of the liquid, and is called the <u>equilibrium vapour pressure</u>.

The phenomenon of vapour pressure is interpreted in terms of molecules of liquid escaping into the empty space above the liquid. In order for the molecules to escape from the liquid phase into the vapour phase, the intermolecular forces (in order of increasing strength: Van der Waals, dipole-dipole, hydrogen bonding) have to be overcome which requires energy. Since the nature of the intermolecular forces is determined by the molecular structure, then the amount of energy required to vapourise the sample also depends on the molecular structure an example of the relationship between structure and properties. As the number of molecules in the vapour above the liquid becomes larger, the rate of return of the molecules from the vapour to the liquid increases until the rate of return is



equal to the rate of escape. This is the equilibrium condition and the corresponding concentration of molecules in the vapour space gives rise to the equilibrium vapour pressure At higher temperatures, the greater average kinetic energy of the molecules in the liquid results in a greater constant rate of escape. Equilibrium is established at higher temperatures, and so larger numbers of molecules are present in the vapour phase and the pressure is higher

When the vapour pressure of a liquid is equal to the atmospheric (or applied) pressure then boiling occurs. The temperature at which this occurs, for a given pressure, is the <u>boiling point</u>. It should be noted, therefore, that the boiling point of a liquid *decreases* as the atmospheric (or applied) pressure *decreases*. This is illustrated by the Vapour Pressure-Temperature Curve above. For example, at sea level the atmospheric pressure is 760 mm Hg (also expressed as 760 torr, 101325 Pa, 101.3 kPa, 1013.25 mbar or 14.696 psi) and pure water boils at 100°C. However, in Calgary (approx. 1050m above sea level) the atmospheric pressure is approximately 670 mm Hg, and water boils at about 96.6°C.

As a rule of thumb, the boiling point of many liquids will drop about 0.5°C for a 10 mm decrease in pressure in the vicinity of 760 mm Hg. At lower pressures, a 10° drop in boiling point is observed for each halving of the pressure. An *approximate measure* that seems to work relatively well in Calgary for calculating the boiling point of a liquid in Calgary is to subtract 1° for every 15° of temperature above 50°C.

So, in order to convert an experimental measurement taken in Calgary (which is at higher altitude than sea level and so is at a lower pressure) to that reported for sea level (higher pressure) one needs to **ADD** a correction factor since the boiling point at sea level is higher than that at higher altitudes.

A more scientific method for correcting boiling point requires knowing the atmospheric pressure (in mmHg), P_{obs}, when and where the boiling point, BP_{obs} is measured:

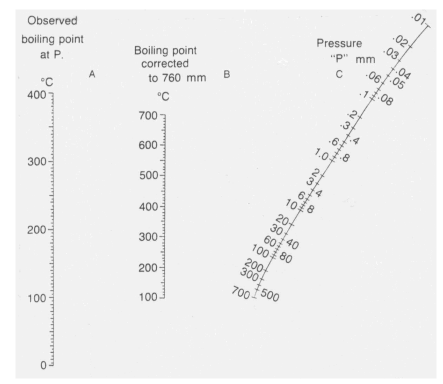
$$BP_{corr} = BP_{obs} - (P_{obs} - 760 \text{mmHg}) \times 0.045 \text{ }^{\circ}\text{C/mmHg}$$

This is the method you should use for correcting boiling points. As an example, the boiling point of water at the summit of Mt Temple (3544m, near Lake Louise, summit pressure approximately 500 mmHg), based on 100 °C at sea level would be:

 $100 \ ^{\circ}C = BP_{obs} - (500 \ mmHg - 760 mmHg) \times 0.045 \ ^{\circ}C/mmHg$ $100 \ ^{\circ}C = BP_{obs} - (-260 mmHg) \times 0.045 \ ^{\circ}C/mmHg = BP_{obs} + 11.7 \ ^{\circ}C/mmHg$ $BP_{obs} = 88.3 \ ^{\circ}C$

In order to use this better method for correcting boiling point, one needs to know the atmospheric pressure at the time when the boiling point was measured. Here is the link to the Environment Canada website for that information. This provides you with the required pressure data at the time pf your laboratory session. Note that 1 kP = 7.50062 mmHg

http://calgary.weatherstats.ca/charts/pressure_station-24hrs.html



At lower pressures, a boiling point nomograph or temperature-pressure alignment chart (shown below) can be used.

A pressure nomograph, used to correct boiling points

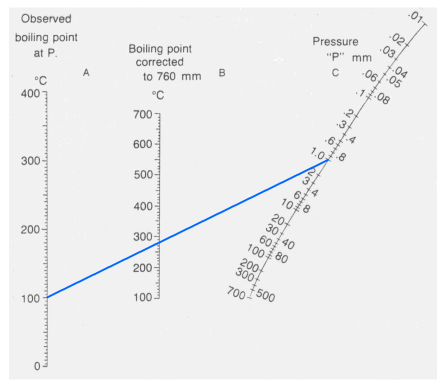
How to use the pressure nomograph.

The basic principle is that a line through two known points on any two different scales (A,B,C) can be used to read off the value on the third scale.

There are two ways in which a pressure nomograph can be used (i) to determine the boiling point at atmospheric pressure (760 mm Hg) given the boiling point at a lower pressure and (ii) to determine the boiling point at a lower pressure given the boiling point at atmospheric pressure.

First, let's say we have a compound with a boiling point of 100°C at 1mm Hg pressure. What is the boiling point at 760mm Hg?

To do this we need to draw a line from 100°C on scale A (left side, observed boiling point) to 1.0 mm Hg on scale C (right side, pressure "P" mm). We can then read off the boiling point at 760 mm on line B, it is about 280°C.

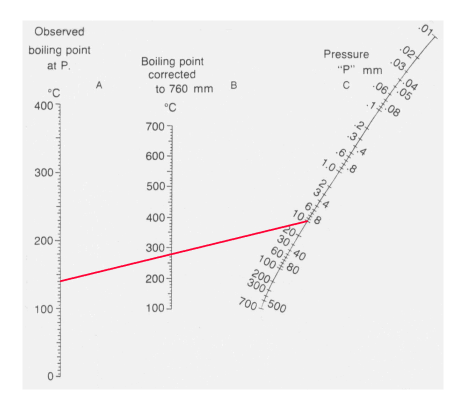


Using a pressure nomograph:

Determining the boiling point at 760mm Hg (scale B) for a sample that boils at 100°C (scale A) at 1mm Hg (scale C)

Now what temperature would that same compound boil at 10 mm Hg pressure? Now we draw a line that passes through 280°C on scale B (middle scale, the boiling point at 760mm Hg) and to 10mm Hg on scale C. By extending that line to scale A, we can read off the new boiling point on scale A (left side) as being about 140°C.

Of course, you don't really have to "draw" the line, it can be done just using the edge of a ruler or something else straight.



Using a pressure nomograph:

Determining the boiling point (scale A) at 10mm Hg (scale C) for a sample that boils at 280°C at 760mm Hg (scale B)

It is also sometimes necessary to apply a thermometer "stem correction" (due to the small thermal contact area and the difference in the coefficient of expansion of mercury or other liquid in a conventional thermometer and the glass). For more information on the thermometer stem correction, see Pasto and Johnson, "Laboratory Text for Organic Chemistry", Prentice Hall.

Boiling point can be determined in a few different ways. If a reasonably large volume of sample is available, then the boiling point of the liquid can be determined using a simple distillation apparatus setup (see simple distillation, technique T.6). However, when large volumes are not available, alternative experimental procedures have to be used. The micro-boiling point method described below can be carried out with just a few mL of a liquid sample and is a reasonably accurate and convenient way to determine the boiling point.

<u>Micro-Boiling Point Determination</u> (quick link to videos : <u>version 1</u>, <u>version 2</u>)

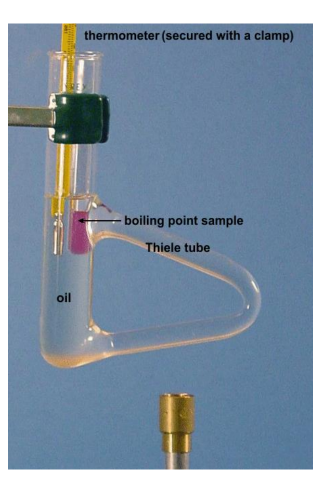


Before using a Bunsen burner make sure all flammable materials (*e.g.* solvents) are removed from the area around the Bunsen burner. This means not only your workspace but also all the students near to you.

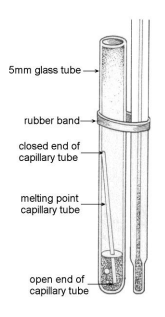
When using the Bunsen burner, make sure that you adjust it to a small flame.

If you are unsure of how to set up and use a Bunsen burner, see the "<u>heat sources</u>" techniques document and the video on the <u>Bunsen burner</u>.

If only small amounts of liquid material are available for boiling point determination, then a micro boiling point apparatus based on a Theile tube should be used. The Thiele tube is a glass tube designed to contain heating oil and a thermometer to which a micro test tube containing the boiling point sample is attached. The shape of the Thiele tube allows for formation of convection currents in the oil when it is heated. These currents maintain a fairly uniform temperature distribution throughout the oil in the tube. The side arm of the tube is designed to generate these convection currents and thus transfer the heat from the flame evenly and rapidly throughout the heating oil. Don't clamp the Thiele tube too tightly otherwise it might crack as it expands as it is heated, yet of course it needs to be secure.



The sample liquid is introduced by Pasteur pipette into a micro test tube (no more than 0.5ml, which is about 10mm depth in the small test tube), and a piece of melting point capillary tubing (sealed at one end) is dropped in with the open end down. The micro test tube assembly is then attached to a thermometer with a rubber band or a thin slice of rubber tubing. The whole unit is then placed in a Thiele tube.



The placement of the test tube / thermometer unit in the Theile tube is important:

- i. the micro test tube should be on the same side of the Thiele tube as the elbow
- **ii.** neither the test tube nor the thermometer bulb should be touching the glass walls of the Thiele tube
- iii. the base of the micro test tube should be just below the joint to the upper part of the elbow (see photograph above)
- iv. the rubber band should be placed well above the level of the oil in the Thiele tube, and,
- v. the oil level should be just above the top of the top elbow joint.
 If the rubber band enters the oil, the band may soften and break in the hot oil allowing the micro test tube to fall into the oil.
 When positioning the band, one should bear in mind that the oil will expand when it is heated.

Once the set up has been complete, the lower part of the side arm of the Thiele tube is **carefully** heated with a **small flame** from the Bunsen burner moving the flame back and forth along the arm. During the heating, there is an <u>initial</u> stream of bubbles as air is expelled and then, a little later, a rapid and continuous stream of bubbles emerges from the inverted capillary tube. At this point stop heating. Soon the stream of bubbles will slow down and stop. When they stop, the liquid sample will be drawn up in to the capillary tube. The moment when the liquid enters the capillary corresponds to the boiling point of the liquid, and the temperature should be recorded.

The explanation of this method is a reasonably simple one. During the initial heating, the air trapped in the capillary tube expands and leaves the tube and vapour from the liquid also enters the tube. There is always vapour in equilibrium with a heated liquid. This gives rise to the initial stream of bubbles. When the temperature reaches the boiling point, the vapour pressure inside the capillary tube equals the atmospheric pressure. As the temperature rises just above the boiling point then the vapour will start to escape : the second set of bubbles. Once the heating is stopped, the only vapour left in the capillary comes from the heated liquid which seals its open end. As the liquid cools, its vapour pressure will decrease and when the vapour pressure drops just below atmospheric pressure, the liquid will be drawn into the capillary tube (forced there by the higher atmospheric pressure).

Two problems are common to this method. The first arises when the liquid sample in the micro test tube is heated so strongly that it evaporates or is boiled away. Once the oil has cooled, add more liquid sample to the small test tube and resume heating but more gently. The second arises when the liquid is not heated above its boiling point. If the heating is stopped at any point below the boiling point of the

liquid, the liquid will enter the tube immediately. It will enter the tube because the trapped vapour will have a pressure less than that of the atmosphere.

Here are links to two very similar videos showing how to carry out a micro boiling point determination, <u>version 1</u>, <u>version 2</u> (newer).

Don't forget to apply a correction due to the reduced atmospheric pressure in Calgary (altitude). To ensure accuracy, you should carry out at least two separate boiling point determinations on your unknown sample.