Skills to Develop

- Compare properties of Group 15 elements.
- Explain the major application of phosphate.
- Describe the equilibria of the ionization of phosphoric acid.

Phosphorus (P) is an essential part of life as we know it. Without the phosphates in biological molecules such as ATP, ADP and DNA, we would not be alive. Phosphorus compounds can also be found in the minerals in our bones and teeth. It is a necessary part of our diet. In fact, we consume it in nearly all of the foods we eat. Phosphorus is quite reactive. This quality of the element makes it an ideal ingredient for matches because it is so flammable. Phosphorus is a vital element for plants and that is why we put phosphates in our fertilizer to help them maximize their growth.

Introduction

Phosphorus plays a big role in our existence but it can also be dangerous. When fertilizers containing phosphorus enter the water, it produces rapid algae growth. This can lead to eutrophication of lakes and rivers; i.e., the ecosystem has an increase of chemical nutrients and this can lead to negative environmental effects. With all the excess phosphorus, plants grow rapidly then die, causing a lack of oxygen in the water and an overall reduction of water quality. It is thus necessary to remove excess phosphorus from our wastewater. The process of removing the phosphorus is done chemically by reacting the phosphorus with compounds such as ferric chloride, ferric sulfate, and aluminum sulfate or aluminum chlorohydrate. Phosphorus, when combined with aluminum or iron, becomes an insoluble salt. The solubility equilibrium constants of \(\text{FePO}_4\) and \(\text{AlPO}_4\) are \(1.3 \times 10^{-22}\) and \(5.8 \times 10^{-19}\), respectively. With solubility's this low, the resulting precipitates can then be filtered out.

Figure 1. Phosphates can lead to excessive algae growth, which can be undesirable

Another example of the dangers of phosphorus is in the production of matches. The flammable nature and cheap manufacturing of white phosphorus made it possible to easily make matches around the turn of the 20th century. However, white phosphorus is highly toxic. Many workers in match factories developed brain damage and a disease called "phosphorus necrosis of the jaw" from exposure to toxic phosphorus vapors. Excess phosphorus accumulation caused their bone tissue to die and rot away. For this reason, we now use red phosphorus or phosphorus sesquisulfide in "safety" matches.
Discovery of Phosphorus

Named from the Greek word phosphoros ("bringer of light"), elemental Phosphorus is not found in its elemental form because this form is quite reactive. Because of this factor it took a long period of time for it to be "discovered". The first recorded isolation of phosphorus was by alchemist Hennig Brand in 1669 involving about 60 pails of urine. After letting a large amount of urine putrefy for a long time, Brand distilled the liquid to a paste, heated the paste, discarded the salt formed, and put the remaining substance under cold water to form solid white phosphorus. Brand's process was not very efficient; the salt he discarded actually contained most of the phosphorus. Nevertheless, he obtained some pure, elemental phosphorus for his efforts. Others of the time improved the efficiency of the process by adding sand, but still continued to discard the salt. Later, phosphorus was manufactured from bone ash. Currently, the process for manufacturing phosphorus does not involve large amounts of putrefied urine or bone ash. Instead, manufacturers use calcium phosphate and coke (Emsley).

Allotropes of Phosphorus

Phosphorus is a nonmetal, solid at room temperature, and a poor conductor of heat and electricity. Phosphorus occurs in at least 10 allotropic forms, the most common (and reactive) of which is so-called white (or yellow) phosphorus which looks like a waxy solid or plastic. It is very reactive and will spontaneously inflame in air so it is stored under water. The other common form of phosphorus is red phosphorus which is much less reactive and is one of the components on the striking surface of a match book. Red phosphorus can be converted to white phosphorus by careful heating.

White phosphorus consists of \(\text{P}_4\) molecules, whereas the crystal structure of red phosphorus has a complicated network of bonding. White phosphorus has to be stored in water to prevent natural combustion, but red phosphorus is stable in air.

When burned, red phosphorus also forms the same oxides as those obtained in the burning of white phosphorus, \(\text{P}_4\text{O}_6\) when air supply is limited, and \(\text{P}_4\text{O}_{10}\) when sufficient air is present.

Diphosphorus (P\(_2\))

Diphosphorus (\(\text{P}_2\)) is the gaseous form of phosphorus that is thermodynamically stable above 1200 °C and until 2000 °C. It can be generated by heating white phosphorus (see below) to 1100 K and is very reactive with a bond-dissociation energy (117 kcal/mol or 490 kJ/mol) half that of dinitrogen (\(\text{N}_2\)).

Figure 2: The four common allotropes of phosphorus. Image used with permission from Wikipedia.
White Phosphorus ($P_4$)

White phosphorus ($P_4$) has a tetrahedral structure. It is soft and waxy, but insoluble in water. Its glow occurs as a result of its vapors slowly being oxidized by the air. It is so thermodynamically unstable that it combusts in air. It was once used in fireworks and the U.S. military still uses it in incendiary bombs.

Figure 3: Structure of white phosphorus. Image used with permission (CC-SA-BY 3.0; Wikipedia)

This [Youtube video](https://www.youtube.com/watch?v=dQw4w9WgXcQ) link shows various experiments with white phosphorus, which help show the physical and chemical properties of it. It also shows white phosphorus combusting with air.

Red Phosphorus and Violet Phosphorus (Polymeric)

Red Phosphorus has more atoms linked together in a network than white phosphorus does, which makes it much more stable. It is not quite as flammable, but given enough energy it still reacts with air. For this reason, we now use red phosphorus in matches.

Figure 4: red phosphorus is in safety matches. Image used with permission (CC-SA-BY 3.0; Wikipedia)

Violet phosphorus is obtained from heating and crystallizing red phosphorus in a certain way. The phosphorus forms pentagonal "tubes".
Black Phosphorus (Polymeric)

Black phosphorus is the most stable form; the atoms are linked together in puckered sheets, like graphite. Because of these structural similarities black phosphorus is also flaky like graphite and possesses other similar properties.

Isotopes of Phosphorus

There are many isotopes of phosphorus, only one of which is stable (\(^{31}\text{P}\)). The rest of the isotopes are radioactive with generally very short half-lives, which vary between a few nanoseconds to a few seconds. Two of the radioactive phosphorus isotopes have longer half-lives. \(^{32}\text{P}\) has a half-life of 14 days and \(^{33}\text{P}\) has a half-life of 25 days. These half-lives are long enough to be useful for analysis and for this reason the isotopes can be used to mark DNA.

\(^{32}\text{P}\) played an important role in the 1952 Hershey-Chase Experiment. In this experiment, Alfred Hershey and Martha Chase used radioactive isotopes of phosphorus and sulfur to determine that DNA was genetic material and not proteins. Sulfur can be found in proteins but not DNA, and phosphorus can be found in DNA but not proteins. This made
Phosphorus and Sulfur effective markers of DNA and protein, respectively. The experiment was set up as follows: Hershey and Chase grew one sample of a virus in the presence of radioactive $^{35}\text{S}$ and another sample of a virus in the presence of $^{32}\text{P}$. Then, they allowed both samples to infect bacteria. They blended the $^{35}\text{S}$ and the $^{32}\text{P}$ samples separately and centrifuged the two samples. Centrifuging separated the genetic material from the non-genetic material. The genetic material penetrated the solid that contained the bacterial cells at the bottom of the tube while the non-genetic material remained in the liquid. By analyzing their radioactive markers, Hershey and Chase found that the $^{32}\text{P}$ remained with the bacteria, and the $^{35}\text{S}$ remained in the supernatant liquid. These results were confirmed by further tests involving the radioactive Phosphorus.

**Phosphorus and Life**

We get most elements from nature in the form of minerals. In nature, phosphorus exists in the form of phosphates. Rocks containing phosphate are fluoroapatite ($\ce{3Ca3(PO4)2.CaF2}$), chloroapatite, ($\ce{3Ca3(PO4)2.CaCl2}$), and hydroxyapatite ($\ce{3Ca3(PO4)2.Ca(OH)2}$). These minerals are very similar to the bones and teeth. The arrangements of atoms and ions of bones and teeth are similar to those of the phosphate containing rocks. In fact, when the $\ce{OH^-}$ ions of the teeth are replaced by $\ce{F^-}$, the teeth resist decay. This discovery led to a series of social and economical issues.

![Diagrams of Ca$_{10}$(PO$_4$)$_2$(OH)$_2$ and Ca$_{10}$(PO$_4$)$_2$F$_2$](image)

**Hydroxyapatite**

**Fluorapatite**

Figure 6: (left) Fluoride ions ($\ce{F^-}$) replace hydroxyl groups ($\ce{OH^-}$) in hydroxyapatite to form fluorapatite in the tooth enamel. (right) A portion of the apatite crystal lattice is depicted showing the replacement of hydroxide for fluoride (big blue circles). Image used with permission (Public Domain; Delmar Larsen).

Nitrogen, phosphorus and potassium are key ingredients for plants, and their contents are key in all forms of fertilizers. From an industrial and economical view point, phosphorus-containing compounds are important commodities. Thus, chemistry of phosphorus has academic, commercial and industrial interests.
As a member of the Nitrogen Family, Group 15 on the Periodic Table, Phosphorus has 5 valence shell electrons available for bonding. Its valence shell configuration is $3s^23p^3$. Phosphorus forms mostly covalent bonds. Any phosphorus rock can be used for the production of elemental phosphorus. Crushed phosphate rocks and sand ($\ce{SiO2}$) react at 1700 K to give phosphorus oxide, $\ce{P4O10}$:

\[
\ce{2 Ca3(PO4)2 + 6 SiO2 \rightarrow P4O10 + 6 CaSiO3} \quad \text{(1)}
\]

$\ce{P4O10}$ can be reduced by carbon:

\[
\ce{P4O10 + 10 C \rightarrow P4 + 10 CO} \quad \text{(2)}
\]

Waxy solids of white phosphorus are molecular crystals consisting of $\ce{P4}$ molecules. They have an interesting property in that they undergo spontaneous combustion in air:

\[
\ce{P4 + 5 O2 \rightarrow P4O10} \quad \text{(3)}
\]

The structure of $\ce{P4}$ can be understood by thinking of electronic configuration ($s^2p^3$) of $\ce{P}$ in bond formation. Sharing three electrons with other $\ce{P}$ atoms gives rise to the 6 $\ce{P-P}$ bonds, leaving a lone pair occupying the 4th position in a distorted tetrahedron.

When burned with insufficient oxygen, $\ce{P4O6}$ is formed:

\[
\ce{P4 + 3 O2 \rightarrow P4O6} \quad \text{(4)}
\]

To each of the $\ce{P-P}$ bonds, an $\ce{O}$ atom is inserted.

Burning phosphorus with excess oxygen results in the formation of $\ce{P4O10}$. An additional $\ce{O}$ atom is attached to the $\ce{P}$ directly:

\[
\ce{P4 + 5 O2 \rightarrow P4O10} \quad \text{(5)}
\]

Thus, the oxides $\ce{P4O6}$ and $\ce{P4O10}$ share interesting features. Oxides of phosphorus, $\ce{P4O10}$, dissolve in water to give phosphoric acid,

\[
\ce{P4O10 + 6 H2O \rightarrow 4 H3PO4} \quad \text{(6)}
\]

Phosphoric acid is a polyprotic acid, and it ionizes at three stages:

\[
\ce{H3PO4 \rightarrow H+ + H2PO4-} \quad \text{(7a)}
\]

\[
\ce{H2PO4- \rightarrow H+ + HPO4^2-} \quad \text{(7b)}
\]

\[
\ce{HPO4^2- \rightarrow H+ + PO4^3-} \quad \text{(7c)}
\]
Phosphoric Acid

Phosphoric Acid is a polyprotic acid, which makes it an ideal buffer. It gets harder and harder to separate the hydrogen from the phosphate making the pKₐ values increase in basicity: 2.12, 7.21, and 12.67. The conjugate bases H₂PO₄⁻, HPO₄²⁻, and PO₄³⁻ can be mixed to form buffer solutions.

Table 1: Ionization constants for the successive deprotonation of phosphoric acid states

<table>
<thead>
<tr>
<th>Reaction</th>
<th>Dissociation Constant</th>
</tr>
</thead>
<tbody>
<tr>
<td>(H_3PO_4 + H_2O \rightarrow H_3O^+ + H_2PO^{4-})</td>
<td>(K_{a1} = 7.5 \times 10^{-3})</td>
</tr>
<tr>
<td>(H_2PO^{4-} + H_2O \rightarrow H_3O^+ + HPO_4^{2-})</td>
<td>(K_{a2} = 6.2 \times 10^{-8})</td>
</tr>
<tr>
<td>(H_2PO^{4-} + H_2O \rightarrow H_3O^+ + PO_4^{3-})</td>
<td>(K_{a3} = 2.14 \times 10^{-3})</td>
</tr>
<tr>
<td>Overall: (H_3PO_4 + 3H_2O \rightarrow 3 H_3O^+ + PO_4^{3-})</td>
<td></td>
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</tbody>
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Past and Present Uses of Phosphorus

Commercially, phosphorus compounds are used in the manufacture of phosphoric acid (\(H_3PO_4\)) (found in soft drinks and used in fertilizer compounding). Other compounds find applications in fireworks and, of course, phoshorescent compounds which glow in the dark. Phosphorus compounds are currently used in foods, toothpaste, baking soda, matches, pesticides, nerve gases, and fertilizers. Phosphoric acid is not only used in buffer solutions it is also a key ingredient of Coca Cola and other sodas! Phosphorus compounds were once used in detergents as a water softener until they raised concerns about pollution and eutrophication. Pure phosphorus was once prescribed as a medicine and an aphrodisiac until doctors realized it was poisonous (Emsley).

References

Edit section


Questions

1. About 85% of the total industrial output of phosphoric acid is used
a. in the detergent industry
b. to produce buffer solutions
c. in the paint industry
d. to produce superphosphate fertilizers
e. in the manufacture of plastics

2. What is the product when phosphorus pentoxide $\ce{P4O10}$ reacts with water? Give the formula of the product.

3. What is the phosphorus-containing product when $\ce{PCl3}$ reacts with water? Give the formula.

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### Solutions

1. *Answer... d*

   The middle number, (for example, 6-5-8) specifies the percentage of phosphorus compound in a fertilizer. Phosphorus is an important element for plant life.

2. *Answer* $\ce{H3PO4}$
   
   $\ce{P4O10 + 6 H2O \rightarrow 4 H3PO4}$

3. *Answer* $\ce{H3PO3}$
   
   $\ce{PCl3 + 3 H2O \rightarrow H3PO3 + 3 HCl}$

   This is a weaker acid than $\ce{H3PO4}$.

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### Contributors

#### Edit section

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