Trihydrides of the Group 15 Elements

All five of the Group 15 elements form hydrides of the formula EH₃. Table 1 lists the IUPAC names along with those in more common usage.

Table 1: Traditional and IUPAC (International Union of Pure and Applied Chemistry) names for the Group 15 hydrides.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Traditional name</th>
<th>IUPAC name</th>
</tr>
</thead>
<tbody>
<tr>
<td>NH₃</td>
<td>Ammonia</td>
<td>Azane</td>
</tr>
<tr>
<td>PH₃</td>
<td>Phosphine</td>
<td>Phosphane</td>
</tr>
<tr>
<td>AsH₃</td>
<td>Arsine</td>
<td>Arsane</td>
</tr>
<tr>
<td>SbH₃</td>
<td>Stibine</td>
<td>Stibane</td>
</tr>
<tr>
<td>BiH₃</td>
<td>Bismuthine</td>
<td>Bismuthane</td>
</tr>
</tbody>
</table>

The boiling point and melting point increase increases going down the Group (Table 2) with increased molecular mass, with the exception of NH₃ whose anomalously high melting and boiling points (Figure 1) are a consequence of strong N-H···H hydrogen bonding. A similar (and stronger) effect is observed for the Group 16 hydrides (H₂E).

Table 2: Selected physical properties of Group 15 hydrides.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Mp (°C)</th>
<th>Bp (°C)</th>
<th>ΔHᵣ (kJ/mol)</th>
<th>E-H bond energy (kJ/mol)</th>
<th>H-E-H bond angle (°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NH₃</td>
<td>-77.7</td>
<td>-33.35</td>
<td>-46.2</td>
<td>391</td>
<td>107</td>
</tr>
<tr>
<td>PH₃</td>
<td>-133</td>
<td>-87.7</td>
<td>9.3</td>
<td>322</td>
<td>93.5</td>
</tr>
<tr>
<td>AsH₃</td>
<td>-116.3</td>
<td>-55</td>
<td>172.2</td>
<td>247</td>
<td>92</td>
</tr>
<tr>
<td>SbH₃</td>
<td>-88</td>
<td>-17.1</td>
<td>142.8</td>
<td>255</td>
<td>91.5</td>
</tr>
</tbody>
</table>
The E-H bond strengths decrease down the group and this correlates with the overall stability of each compound (Table 2). The H-E-H bond angles (Table 2) also decrease down the Group. The H-E-H bond angle is expected to be a tetrahedral ideal of 109.5°, but since lone pairs repel more than bonding pairs, the actual angle would be expected to be slightly smaller. Two possible explanations are possible for the difference between NH₃ and the other hydrides.

1. The N-H bond is short (1.015 Å) compared to the heavier analogs, and nitrogen is more electronegative than hydrogen, so the bonding pair will reside closer to the central atom and the bonding pairs will repel each other opening the H-N-H angle more than observed for PH₃, etc.

2. The accessibility of the 2s and 2p orbitals on nitrogen allows for hybridization and the orbitals associated with N-H bonding in NH₃ are therefore close to sp³ in character, resulting in a close to tetrahedral geometry. In contrast, hybridization of the ns and np orbitals for P, As, etc., is less accessible, and as a consequence the orbitals associated with P-H bonding in PH₃ are closer to p in character resulting in almost 90° H-P-H angle. The lower down the Group the central atom the less hybridization that occurs and the closer to pure p-character the orbitals on E associated with the E-H bond.

Ammonia (NH₃) is a colorless, pungent gas (bp = -33.5 °C) whose odor can be detected at concentrations as low 20 – 50 ppm. Its high boiling point relative to its heavier congeners is indicative of the formation of strong hydrogen bonding. The strong hydrogen bonding also results in a high heat of vaporization (23.35 kJ/mol) and thus ammonia can be conveniently used as a liquid at room temperature despite its low boiling point.

WARNING

Ammonia solution causes burns and irritation to the eyes and skin. The vapor causes severe irritation to the respiratory system. If swallowed the solution causes severe internal damage.
Synthesis

Ammonia is manufactured on the industrial scale by the Haber process using the direct reaction of nitrogen with hydrogen at high pressure (10^2 – 10^3 atm) and high temperature (400 – 550 °C) over a catalyst (e.g., α-iron), (8.3.1).

\[
\text{N}_2 + 3 \text{H}_2 \rightarrow 2 \text{NH}_3
\]

On the smaller scale ammonia is prepared by the reaction of an ammonium salt with a base, (8.3.2), or hydrolysis of a nitride, (8.3.3). The latter is a convenient route to ND\textsubscript{3} by the use of D\textsubscript{2}O.

\[
\text{NH}_4^+ \text{X} + \text{OH}^- \rightarrow \text{NH}_3 + \text{H}_2\text{O} + \text{X}^-
\]

\[
\text{Mg}_3\text{N}_2 + \text{H}_2\text{O} \rightarrow 3 \text{Mg(OH)}_2 + 2 \text{NH}_3
\]

Structure

The nitrogen in ammonia adopts sp\textsuperscript{3} hybridization, and ammonia has an umbrella structure (Figure 2) due to the stereochemically active lone pair.

![Figure 2: The structure of ammonia.](image)

In a similar manner to water, (8.3.4), ammonia is a self-ionizing, (8.3.5); however, the equilibrium constant (K = 10^-33) is much lower than water (K = 10^-14). The lower dielectric constant of ammonia (16.5 @ 20 °C) as compared to water (80.4 @ 20 °C) means that ammonia is not as good as water as a solvent for ionic compounds, but is better for covalent organic compounds.

\[
2 \text{H}_2\text{O} \rightleftharpoons \text{H}_3\text{O}^+ + \text{OH}^-
\]

\[
2 \text{NH}_3 \rightleftharpoons \underset{\text{ammonium}}{\text{NH}_4^+} + \underset{\text{amide}}{\text{NH}_2^-}
\]

Reactions

The similarity of ammonia and water means that the two compounds are miscible. In fact, ammonia forms a series of
solid hydrates, analogous to ice in which hydrogen bonding defines the structures (Figure 3). Several hydrates of ammonia are known, including: NH$_3$.2H$_2$O (ammonia dihydrate, ADH), NH$_3$.H$_2$O (ammonia monohydrate, AMH), and 2NH$_3$.2H$_2$O (ammonia hemihydrate, AHH).

![Figure 3: The crystal structure of ammonia monohydrate (ANH-II) with hydrogen atoms omitted. Adapted from A. D. Fortes, E. Suard, M. -H. Lemée-Cailleau, C. J. Pickard, and R. J. Needs, J. Am. Chem. Soc., 2009, 131, 13508. Copyright: American Chemical Society (2009).](image)

It should be noted that these hydrates do not contain discrete NH$_4^+$ or OH$^-$ ions, indicating that ammonium hydroxide does not exist as a discrete species despite the common usage of the name. In aqueous solution, ammonia is a weak base ($pK_b$ = 4.75), (8.3.6).

\[
\text{NH}_3(\text{aq}) + \text{H}_2\text{O}(\text{l}) \rightleftharpoons \text{NH}_4(\text{aq})^+ + \text{OH}^-_\text{(aq)}
\]

Note

Ammonia solutions commonly used in the laboratory is a 35% solution in water. In warm weather the solution develops pressure and the cap must be released with care. The 25% solution sold commercially (for home use) is free from this problem.

Ammonia is a Lewis base and readily forms Lewis acid-base complexes with both transition metals, (8.3.7), and main group metals (Figure 4).

\[
\text{[Ni(H}_2\text{O})_6]^{2+} + 6 \text{NH}_3 \rightleftharpoons \text{[Ni(NH}_3)_6]^{2+} + 6 \text{H}_2\text{O}
\]
The formation of stable ammonia complexes is the basis of a simple but effective method of detection: Nessler’s reagent, (8.3.8). Using a 0.09 mol/L solution of potassium tetraiodomercurate(II), K₂[HgI₄], in 2.5 mol/L potassium hydroxide. A yellow coloration indicates the presence of ammonia: at higher concentrations, a brown precipitate may form. The sensitivity as a spot test is about 0.3 μg NH₃ in 2 μL.

\[\text{NH}_4^+ + 2 [\text{HgI}_4]^{2-} + 4 \text{OH}^- \rightarrow \text{HgO} \cdot \text{Hg(NH}_2\text{I} + 7 \text{I}^- + 3 \text{H}_2\text{O}\]

Ammonia forms a blue solution with Group 1 metals. As an example, the dissolution of sodium in liquid ammonia results in the formation of solvated Na⁺ cations and electrons, (8.3.9) where solv = NH₃. The solvated electrons are stable in liquid ammonia and form a complex: \([\text{e}^-(\text{NH}_3)s]\).

\[\text{Na}_{(s)} \rightleftharpoons \text{Na}^+_{(solv)} + \text{e}^-_{(solv)}\]

It is this solvated electron that gives the strong reducing properties of the solution as well as the characteristic signal in the ESR spectrum associated with a single unpaired electron. The blue color of the solution is often ascribed to these solvated electrons; however, their absorption is in the far infra-red region of the spectrum. A second species, Na⁻(solv), is actually responsible for the blue color of the solution.

\[2 \text{Na}^-_{(solv)} \rightleftharpoons \text{Na}^+_{(solv)} + \text{Na}^-_{(solv)}\]

The reaction of ammonia with oxygen is highly favored, (8.3.11), and the flammability limit of ammonia is 16 – 25 vol%. If the reaction is carried out in the presence of a catalyst (Pt or Pd) the reaction can be limited to the formation of nitric
oxide (NO), (8.3.12).

\[ \text{4 NH}_3 + 3 \text{ O}_2 \rightarrow 2 \text{ N}_2 + 6 \text{ H}_2\text{O} \]

\[ \text{4 NH}_3 + 5 \text{ O}_2 \rightarrow 4 \text{ NO} + 6 \text{ H}_2\text{O} \]

---

**Ammonium salts**

The ammonium cation (NH\(_4^+\)) behaves in a similar manner to the Group 1 metal ions. The solubility and structure of ammonium salts particularly resembles those of potassium and rubidium because of their relative size (Table 3). One difference is that ammonium salts often decompose upon heating, (8.3.13).

\[ \text{NH}_4\text{Cl}_{(s)} \rightarrow \text{NH}_3_{(g)} + \text{HCl}_{(g)} \]

<table>
<thead>
<tr>
<th>Cation</th>
<th>Ionic radius (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>K(^+)</td>
<td>1.33</td>
</tr>
<tr>
<td>NH(_4^+)</td>
<td>1.43</td>
</tr>
<tr>
<td>Rb(^+)</td>
<td>1.47</td>
</tr>
</tbody>
</table>

Table 3: Ionic radius of the ammonium ion compared to those of potassium and rubidium.

The decomposition of ammonium salts of oxidizing acids can often be violent to highly explosive, and they should be treated with care. For example, while ammonium dichromate, (NH\(_4\))\(_2\)Cr\(_2\)O\(_7\), decomposes to give a volcano (Figure 5), ammonium permanganate, NH\(_4\)[MnO\(_4\)], is friction sensitive and explodes at 60 °C. Ammonium nitrate, NH\(_4\)[NO\(_3\)], can cause fire if contacted with a combustible material and is a common ingredient in explosives since it acts as the oxygen source due to its positive oxygen balance, i.e., the compound liberates oxygen surplus to its own needs upon decomposition, (8.3.14).

\[ \text{NH}_4[\text{NO}_3] \rightarrow \text{H}_2\text{O} + \text{N}_2 + \text{O}_2 \]
Ammonia has a reasonable liquid range (-77 to –33 °C), and as such it can be readily liquefied with dry ice (solid CO$_2$, $T_{sub} = -78.5$ °C), and handled in a thermos flask. Ammonia’s high boiling point relative to its heavier congeners is indicative of the formation of strong hydrogen bonding, which also results in a high heat of vaporization (23.35 kJ/mol). As a consequence ammonia can be conveniently used as a liquid at room temperature despite its low boiling point.

Liquid ammonia is a good solvent for organic molecules (e.g., esters, amines, benzene, and alcohols). It is a better solvent for organic compounds than water, but a worse solvent for inorganic compounds. The solubility of inorganic salts is highly dependant on the identity of the counter ion (Table 4).

**Table 4: General solubility of inorganic salts in liquid ammonia as a function of the counter ion.**

<table>
<thead>
<tr>
<th>Soluble in liquid NH$_3$</th>
<th>Generally insoluble in liquid NH$_3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>SCN$^-$, I$^-$, NH$_4^+$, NO$_3^-$, NO$_2^-$, ClO$_4^-$</td>
<td>F$^-$, Cl$^-$, Br$^-$, CO$_3^{2-}$, SO$_4^{2-}$, O$_2^-$, OH$^-$, S$^{2-}$</td>
</tr>
</tbody>
</table>

The difference in solubility of inorganic salts in ammonia as compared to water, as well as the lower temperature of liquid ammonia, can be used to good advantage in the isolation of unstable compounds. For example, the attempted synthesis of ammonium nitrate by the reaction of sodium nitrate and ammonium chloride in water results in the formation of nitrogen and water due to the decomposition of the nitrate, (8.3.15). By contrast, if the reaction is carried out in liquid...
ammonia, the sodium chloride side product is insoluble and the ammonium nitrate may be isolated as a white solid after filtration and evaporation below its decomposition temperature of 0 °C, (8.3.16).

\[
\text{NaNO}_2 + \text{NH}_4\text{Cl} \xrightarrow{\text{H}_2\text{O}} \text{NaCl} + \text{NH}_4(\text{NO}_2) \rightarrow \text{N}_2 + 2 \text{H}_2\text{O}
\]

\[
\text{NaNO}_2 + \text{NH}_4\text{Cl} \xrightarrow{\text{NH}_3} \text{NaCl} \downarrow + \text{NH}_4(\text{NO}_2)
\]

**Ammonation**

Ammonation is defined as a reaction in which ammonia is added to other molecules or ions by covalent bond formation utilizing the unshared pair of electrons on the nitrogen atom, or through ion-dipole electrostatic interactions. In simple terms the resulting *ammine* complex is formed when the ammonia is acting as a Lewis base to a Lewis acid, (8.3.17) and (8.3.18), or as a ligand to a cation, e.g., [Pt(NH3)4]^{2+}, [Ni(NH3)6]^{2+}, [Cr(NH3)6]^{3+}, and [Co(NH3)6]^{3+}.

\[
\text{SiF}_4 + 2 \text{NH}_3 \rightarrow \text{SiF}_4(\text{NH}_3)_2
\]

\[
\text{BF}_3 + \text{NH}_3 \rightarrow \text{BF}_3(\text{NH}_3)
\]

**Ammonolysis**

Ammonolysis with ammonia is an analogous reaction to hydrolysis with water, i.e., a dissociation reaction of the ammonia molecule producing H^+ and an NH_2^- species. Ammonolysis reactions occur with inorganic halides, (8.3.18) and (8.3.19), and organometallic compounds, (8.3.20). In both case the NH_2^- moiety forms a substituent or ligand.

\[
\text{P(O)Cl}_3 + 6 \text{NH}_3 \rightarrow \text{P(O)(NH}_2)_3 + 3 \text{NH}_4\text{Cl}
\]

\[
\text{BCl}_3 + 6 \text{NH}_3 \rightarrow \text{B(NH}_2)_3 + 3 \text{NH}_4\text{Cl}
\]

\[
\text{Al(CH}_3)_3 + \text{NH}_3 \rightarrow \frac{1}{n}[(\text{H}_3\text{C})_2\text{Al(NH}_2)_3]_n + \text{CH}_4
\]

The reaction of esters, (8.3.21), and aryl halides, (8.3.22), are also examples of ammonolysis reactions.

\[
\text{RC(O)OR'} + \text{NH}_3 \rightarrow \text{RC(O)NH}_2 + \text{R'OH}
\]

\[
\text{C}_6\text{H}_5\text{Cl} + 2 \text{NH}_3 \rightarrow \text{C}_6\text{H}_5\text{NH}_2 + \text{NH}_4\text{Cl}
\]

**Homoleptic amides**

A homoleptic compound is a *compound with all the ligands being identical*, e.g., M(NH2)_n. A general route to homoleptic amide compounds is accomplished by the reaction of a salt of the desired metal that is soluble in liquid ammonia (Table 4) with a soluble Group 1 amide. The solubility of the Group 1 amides is given in Table 5. Since all amides are insoluble (except those of the Group 1 metals) are insoluble in liquid ammonia, the resulting amide may be readily isolated, e.g., (8.3.23) and (8.3.24).

\[
\text{Mn(SCN)}_2 + 2 \text{KNH}_2 \rightarrow \text{Mn(NH}_2)_2 \downarrow + 2 \text{KSCN}
\]
\[ \text{Cr(NO}_3\text{)}_3 + 3 \text{KNH}_2 \rightarrow \text{Cr(NH}_2\text{)}_3 \downarrow + 3 \text{KNO}_3\]

Table 5: Solubility of Group amides in liquid ammonia.

<table>
<thead>
<tr>
<th>Amide</th>
<th>Solubility in liquid ammonia</th>
</tr>
</thead>
<tbody>
<tr>
<td>LiNH(_2)</td>
<td>Sparingly soluble</td>
</tr>
<tr>
<td>NaNH(_2)</td>
<td>Sparingly soluble</td>
</tr>
<tr>
<td>KNH(_2)</td>
<td>Soluble</td>
</tr>
<tr>
<td>RbNH(_2)</td>
<td>Soluble</td>
</tr>
<tr>
<td>CsNH(_2)</td>
<td>Soluble</td>
</tr>
</tbody>
</table>

**Redox reactions**

Ammonia is poor as an oxidant since it is relatively easily oxidized, e.g., (8.3.25) and (8.3.26). Thus, if it is necessary to perform an oxidation reaction ammonia is not a suitable solvent; however, it is a good solvent for reduction reactions.

\[ 4 \text{NH}_3 + 5 \text{O}_2 \rightarrow 4 \text{NO} + 6 \text{H}_2\text{O} \]

\[ 2 \text{NH}_3 + 3 \text{CuO} \rightarrow \text{N}_2 + 3 \text{H}_2\text{O} + 3 \text{Cu} \]

Liquid ammonia will dissolve Group 1 (alkali) metals and other electropositive metals such as calcium, strontium, barium, magnesium, aluminum, europium, and ytterbium. At low concentrations (ca. 0.06 mol/L), deep blue solutions are formed: these contain metal cations and solvated electrons, (8.3.27). The solvated electrons are stable in liquid ammonia and form a complex: \([e^-(\text{NH}_3)_6]\).

\[ \text{Na}_\text{(s)} \rightarrow \text{Na}_\text{(solv)} \rightleftharpoons \text{Na}^+_{\text{(solv)}} + e^-_{\text{(solv)}} \]

The solvated electrons provide a suitable and powerful reducing agent for a range of reactions that are not ordinarily accomplished, e.g., (8.3.28) and (8.3.29).

\[ [\text{Ni(CN)}_4]^{2-} + 2 e^-_{\text{(solv)}} \rightarrow [\text{Ni(CN)}_4]^{4-} \]

\[ [\text{Co}_2(\text{CO})_8] + 2 e^-_{\text{(solv)}} \rightarrow 2 [\text{Co(CO)}_4]^- \]

**Ammonia: From Uses in Agriculture and Beer Production to Prolonging the First World War**

One of the cornerstones of the industrial expansion in Germany between 1870 and 1910 (when the population grew from 41 million to 66 million) was the chemicals industry. In particular companies such as Badische Anilin und Soda Fabrik (Baden Aniline and Soda Factory or BASF) had developed their business on the formation of synthetic dyes derived from the chemicals in coal tar.
Coal tar was formed as a by-product of the gasification of coal. A key ingredient of coal tar is naphtha (Table 6) that was used by the rubber industry; however, the remainder was disposed off. It was not long before chemists started to investigate the constituents present in coal tar. One of the first discoveries was by a chemistry student at the Royal College of Chemistry (now Imperial College) in London, William Perkin (Figure 6), who isolated the first synthetic dye: mauveine (Figure 7). This discovery led to an explosion in the chemical industry based on the extraction of compounds from coal tar, which was an essentially free waste product from every gasworks. Nowhere was this research more commercially driven that in Germany.

Table 6: The typical physical properties of naphtha.

<table>
<thead>
<tr>
<th>Property</th>
<th>Typical value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Molecular weight</td>
<td>100 - 215 g/mol</td>
</tr>
<tr>
<td>Specific gravity</td>
<td>0.75 - 0.85 g/cm³</td>
</tr>
<tr>
<td>Boiling point</td>
<td>160 - 220 °C</td>
</tr>
<tr>
<td>Vapor pressure</td>
<td>5 mmHg</td>
</tr>
<tr>
<td>Solubility in water</td>
<td>Insoluble</td>
</tr>
</tbody>
</table>

Figure 6: A portrait of English chemist Sir William Perkin (1838 - 1907) holding a sample of cloth dyed with his discovery, mauveine, which is often called Perkin’s purple.
The rapid increase in the German population also put a strain on the countries food resources. What compounded this issue was that the aristocratic Junkers families of East Prussia who owned much of the land in what was known as Germany’s breadbasket. Junkers grew rye on their estates because the soil was too light for wheat, and since rye was fertilized with potash (potassium oxide, K$_2$O) of which Germany had vast resources. However, in 1870 grain from the US was becoming cheaper and thus competitive with German rye. To protect their profits the Junkers demanded both subsidies for the export of rye and tariffs for the import of wheat. The result of this was that all the German rye was leaving the country and there was not enough wheat being produced to satisfy the needs of the local population. Sufficient wheat could be grown in Germany if a suitable fertilizer was available.

Sodium nitrate (NaNO$_3$), also known as Chile saltpeter, was the most common fertilizer. Unfortunately, by 1900 the deposits looked to be depleted and an alternative was needed. The alternative was found as a component in coal tar. It was known that one of the chemicals that caused the stink associated with coal gas and coal tar was ammonia (NH$_3$). Chemist George Fownes (Figure 8) had suggested that ammonia be turned into a salt and used as a fertilizer. Unfortunately, the amount of ammonia that could be separated from coal tar was still insufficient, so if ammonia could be made on a large enough scale, then large-scale manufacture of a fertilizer could be realized.
In 1909 Fritz Haber (Figure 9) presented a method of ammonia synthesis to BASF. His work in collaboration with Carl Bosch (Figure 10) resulted in the process known as the Haber-Bosch process in which nitrogen and hydrogen are mixed at high temperature (600 °C) under pressure (200 atm) over an osmium catalyst, (8.3.30).

\[
\text{N}_2 + 3 \text{H}_2 \xrightarrow{\text{Os cat}} 6 \text{NH}_3
\]
Figure 9: German chemists Fritz Haber (1868 –1934) who received the Nobel Prize in Chemistry in 1918 for his development for synthesizing ammonia.
German chemist Carl Bosch (1874 –1940) who received the Nobel Prize in Chemistry in 1931 for his work in high-pressure chemistry.

It is interesting to note that the realization of the Haber-Bosch process required not only high-pressure vessels to be constructed by the steel industry, but also the liquid forms of nitrogen and hydrogen. As it turned out ammonia was a necessary component for enabling the production of liquid nitrogen and hydrogen, and involved a false hypothesis of what caused malaria, which led to a desire to keep drinks cold.

Long before it was understood the real cause of malaria, John Gorrie (Figure 11), a doctor working in Apalachicola on the Gulf Coast of Florida, was obsessed with finding a cure for malaria. The term malaria originated from Medieval Italian: mala aria (bad air), and it was associated with swamps and marshlands. Gorrie noticed that malaria was connected to hot humid weather so he began hanging bowl of ice in wards and circulating the air with a fan. However, ice was cut from frozen lakes and rivers in the North East of the US, stored and then shipped all over the world, and Apalachicola was so small that ice was seldom delivered. Gorrie started looking into methods of making ice. It was well known that when a compressed gas expands it takes heat from its surroundings. Gorrie made a steam engine that compressed air in a piston, which when the piston retracted the air cooled. On the next compression stroke the cold air was pushed out across a brine solution (saturated aqueous NaCl) cooling it. When he brought water in contact with the cold brine, it froze creating the first man made ice. On 14 July 1850 Gorrie produced ice for the French Consul to cool the champagne for the celebration of Bastille Day. Just before he died, Gorrie suggested that his (by then) Patented process could be used for cooling food for transport, and it was this application that was used extensively by British merchants to bring meat from Australia to Britain. However, in Germany, Gorrie’s invention was more useful for beer.
Figure 11: Portrait of American physician and scientist John Gorrie (1803 –1855) is considered the father of refrigeration and air conditioning.

Whereas the British traditionally brewed beer in which the yeast ferments on the surface (top fermentation) at a temperature of 60 – 70 °F, in Germany beer was made using bottom fermentation. This style of fermentation requires a temperature just above freezing. Traditionally, cold cellars were used to store the fermenting beer, and it is from here the name lager is derived from the German verb *largern*: to store. There had been a law in Germany preventing brewing in the summer, but with Gorrie’s process the possibility was to be able to brew beer all year. Carl von Linde (Figure 12) was asked to develop a refrigeration system. He used ammonia instead of air in Gorrie’s system, and in 1879 he set up a company to commercialize his ideas. The success of his refrigerator was such that by 1891 there were over 12,000 fridges being used, and more importantly there was now a convenient method of liquefying gases such as hydrogen and nitrogen; both of which were needed for the Haber-Bosch process.
Figure 12: German engineer Carl Paul Gottfried von Linde (1842 - 1934).

As a consequence of the use of ammonia as a refrigerant, it was possible to prepare ammonia on a large industrial scale. Ammonia prepared by the Haber-Bosch process can be converted to nitric acid by the Ostwald process developed by Wilhelm Ostwald (Figure 13). Treatment of ammonia with air over a platinum catalyst yields initially nitric oxide, (8.3.31), and subsequently to nitrogen dioxide, (8.3.32), which dissolves in water to give nitric acid, (8.3.33).

\[
\begin{align*}
4 \text{NH}_3 + 5 \text{O}_2 &\xrightarrow{\text{Pt (cat)}} 4 \text{NO} + 6 \text{H}_2\text{O} \\
2 \text{NO} + \text{O}_2 &\xrightarrow{\text{Pt (cat)}} 2 \text{NO}_2 \\
3 \text{NO}_2 + \text{H}_2\text{O} &\rightarrow 2 \text{HNO}_3 + \text{NO}
\end{align*}
\]

Addition of soda (sodium hydroxide, NaOH) to nitric acid results in the formation of sodium nitrate, (8.3.34), which was the same fertilizer produced from the deposits in Chile.

\[
\begin{align*}
\text{HNO}_3 + \text{NaOH} &\rightarrow \text{NaNO}_3 + \text{H}_2\text{O}
\end{align*}
\]
Figure 13: German chemist Friedrich Wilhelm Ostwald (1853 – 1932) received the Nobel Prize in 1909 for his work on catalysis.

Unfortunately, for the Haber-Bosch and Ostwald processes, an even cheaper form of fertilizer was synthesized around the same time using calcium carbide to prepare calcium cyanamide (CaCN₂), (8.3.35). As a consequence, the Haber-Bosch process was forgotten until the outbreak of the First World War in 1914.

\[
\text{CaC}_2 + \text{N}_2 \rightarrow \text{CaCN}_2 + \text{C}
\]

Within weeks of the outbreak Germany realized it had only enough explosives for about a year of conflict. This was because the main source of explosives, sodium nitrate was the same source that gave fertilizer, i.e., Chile. Realizing this, the Royal Navy effectively blockaded the supply lines. If Germany did not find another source of Great War would have been over early in 1916, however, it was remembered that the Haber-Bosch process in combination with the Ostwald processes would allow the synthesis of nitric acid, which when mixed with cotton, made nitrocellulose (Figure 14), also known as gun cotton, an explosive, (8.3.36).

\[
\text{HNO}_3 + \text{C}_6\text{H}_{10}\text{O}_5 \rightarrow \text{C}_6\text{H}_7(\text{NO}_2)_3\text{O}_5 + 3 \text{H}_2\text{O}
\]

Figure 14: The structure of nitrocellulose.
As a result of the industrial synthesis of ammonia Germany was able to manufacture sufficient explosives to fight until 11 November 1918, by which time almost 10 million were dead, almost 7 million missing, and over 21 million were wounded (Figure 15).

**Figure 15**: The Douamont Ossuary cemetery and World War I memorial in Verdun, France.

---

**Hydrazine**

Hydrazine (N\(_2\)H\(_4\)) is a colorless liquid with an odor similar to ammonia. Hydrazine has physical properties very close to water, with a melting point of 2 °C and a boiling point of 114 °C. The similarity in its chemistry to water is as a result of strong intermolecular hydrogen bonding.

**WARNING**

Hydrazine is highly toxic and dangerously unstable, and is usually handled as aqueous solution for safety reasons. Even so, hydrazine hydrate causes severe burns on the skin and eyes. Contact with transition metals, their oxides (e.g., rust), or salts cause catalytic decomposition and possible ignition of evolved hydrogen. Reactions with oxidants are violent.

**Synthesis**

Hydrazine is manufactured on the industrial scale by the Olin Raschig process using the reaction of a sodium hypochlorite solution with ammonia at 5 °C to form chloramine (NH\(_2\)Cl) and sodium hydroxide, (8.3.37). The chloramines solution is reacted with ammonia under pressure at 130 °C, (8.3.38). Ammonia is used in a 33 fold excess.

\[
\text{NH}_3 + \text{OCl}^- \rightarrow \text{OH}^- + \text{NH}_2\text{Cl}
\]

\[
\text{NH}_2\text{Cl} + \text{OH}^- + \text{NH}_3 \rightarrow \text{N}_2\text{H}_4 + \text{Cl}^- + \text{H}_2\text{O}
\]

If transition metals are present then decomposition occurs, (8.3.38), and therefore, ethylenediaminetetraacetic acid (EDTA, Figure 16) is added to complex the transition metal ions. The as produced hydrazine solution can be concentrated by distillation to give a 65% solution. Anhydrous hydrazine is formed by the distillation from NaOH.

\[
\text{2NH}_2\text{Cl} + \text{N}_2\text{H}_4 \rightarrow 2\text{NH}_4^+ + 2\text{Cl}^- + \text{N}_2
\]
Alternative routes to hydrazine include the oxidation of urea with sodium hypochlorite, (8.3.40), and the reaction of ammonia and hydrogen peroxide, (8.3.41).

\[
\text{H}_2\text{N}_2\text{C}=\text{O} + \text{NaOCl} + 2 \text{NaOH} \rightarrow \text{N}_2\text{H}_4 + \text{H}_2\text{O} + \text{NaCl} + \text{Na}_2\text{CO}_3
\]

\[
2 \text{NH}_3 + \text{H}_2\text{O}_2 \rightarrow \text{N}_2\text{H}_4 + 2 \text{H}_2\text{O}
\]

**Structure**

The nitrogen atoms in hydrazine adopt sp\(^3\) hybridization (Figure 17a), and molecule adopts a gauche conformation in the vapor, liquid and solid states (Figure 17b).

![Figure 17: The structure of hydrazine.](image)

In a similar manner to ammonia, (8.3.42), hydrazine is a self-ionizing, (8.3.43). While there is a wide range of salts of the \(\text{N}_2\text{H}_5^+\) cation, only the sodium and potassium salts of \(\text{N}_2\text{H}_3^-\) are stable.

\[
2 \text{NH}_3 \rightleftharpoons \underset{\text{ammonium}}{\text{NH}_4^+} + \underset{\text{amide}}{\text{NH}_2^-}
\]

\[
2 \text{N}_2\text{H}_4 \rightleftharpoons \text{N}_2\text{H}_5^+ + \text{N}_2\text{H}_3^-
\]

**Reaction chemistry and uses**

Hydrazine is polar, highly ionizing, and forms stable hydrogen bonds, and its resemblance to water is reflected in the formation of aqueous solutions and hydrates. In the solid state the monohydrate is formed, i.e., \(\text{N}_2\text{H}_4.\text{H}_2\text{O}\). In solution
Hydrazine acts as a base to form the hydrazinium ion, (8.3.44) where $K_b = 8.5 \times 10^{-7}$. The presence of a second Lewis base site means that hydrazine can be protonated twice to form the hydrazonium ion, (8.3.45) where $K_b = 8.9 \times 10^{-16}$.

Salts of the cation $N_2H_5^+$ are stable in water; however, the salts of the dication are less stable.

\[
\text{2 NH}_3 \rightleftharpoons \text{NH}_4^+ + \text{NH}_2^- \\
\text{N}_2\text{H}_5^+ + \text{H}_2\text{O} \rightleftharpoons \text{N}_2\text{H}_5^{2+} + \text{OH}^- \\
\]

The reaction of hydrazine with oxygen is highly favored, (8.3.46), and the explosive limit is $1.8 \text{ – } 100 \text{ vol}\%$.

\[
\text{N}_2\text{H}_4 + \text{O}_2 \rightarrow \text{N}_2 + 2 \text{H}_2\text{O} \\
\]

Hydrazine is useful in a number of organic reactions for the synthesis of a wide range of compounds used in pharmaceuticals, textile dyes, and in photography, including:

- Hydrazone formation, (8.3.47) and (8.3.48).
- Alkyl-substituted hydrazine synthesis via direct alkylation with alkyl halides.
- Reaction with 2-cyanopyridines to form amide hydrazides, which can be converted using 1,2-diketones into triazines.
- Use in the Wolff-Kishner reduction that transforms the carbonyl group of a ketone or aldehyde into a methylene (or methyl) group via a hydrazone intermediate, (shown below).
- As a building block for the preparation of many heterocyclic compounds via condensation with a range of difunctional electrophiles.
- Cleavage N-alkylated phthalimide derivatives.
- As a convenient reductant because the by-products are typically nitrogen gas and water.

\[
\text{2 (CH}_3\text{)}_2\text{C=O} + \text{N}_2\text{H}_4 \rightarrow 2 \text{H}_2\text{O} + [\text{(CH}_3\text{)}_2\text{C=N}]_2 \\
[\text{(CH}_3\text{)}_2\text{C=N}]_2 + \text{N}_2\text{H}_4 \rightarrow 2 \text{(CH}_3\text{)+2\text{C=N}NH}_2 \\
\]

**Messerschmitt Me 163 Komet**

Designed by Alexander Lippisch (Figure 18), the Messerschmitt Me 163B Komet (Figure 19) was the first rocket-powered fighter plane. With a top speed of around 596 mph (Mach 0.83) and a service ceiling of 40,000 ft, the Komet’s performance of the Me 163B far exceeded that of contemporary piston engine fighters. However, despite its impressive performance, it was only produced in limited numbers (ca. 370 as compared to the 1,430 built of its jet powered compatriot the Me 262) and was not an effective combat airplane.
The Komet was powered by the HWK 109-509 *hot engine* (Figure 20) that used a mixture of a fuel and an oxidizer. The fuel was a mixture of hydrazine hydrate (30%), methanol (57%), and water (13%) that was designated by the code name, C-Stoff, that burned with the oxygen-rich exhaust from hydrogen peroxide (T-Stoff) used as the oxidizer. The C-Stoff was stored in a glass tank on the plane, while the T-Stoff was stored in an aluminum container. An oxidizing agent cocktail of CaMnO$_4$ and/or K$_2$CrO$_4$ was added to the T-Stoff generating steam and high temperatures, this in turn reacted violently with the C-Stoff. The flow of reagents was controlled by two pumps, to regulate the rate of combustion and thereby the amount of thrust. The violent combustion process resulted in the formation of water, carbon dioxide and nitrogen, and a huge amount of heat sending out a superheated stream of steam, nitrogen and air that was drawn in through the hole in the mantle of the engine, thus providing a forward thrust of approximately 3,800 lbf. Because of the potential hazards of mixing the fuels, they were stored at least $1/2$ mile apart, and the plane was washed with water between fueling steps and after missions.
Phosphine and Arsine

Because of their use in metal organic chemical vapor deposition (MOCVD) of 13-15 (III-V) semiconductor compounds phosphine (PH$_3$) and arsine (AsH$_3$) are prepared on an industrial scale.

**Synthesis**

Phosphine (PH$_3$) is prepared by the reaction of elemental phosphorus (P$_4$) with water, (8.3.49). Ultra pure phosphine that is used by the electronics industry is prepared by the thermal disproportionation of phosphorous acid, (8.3.50).

\[
2 P_4 + 12 H_2O \rightarrow 5 PH_3 + 3 H_3PO_4
\]

\[
4 H_3PO_4 \rightarrow PH_3 + 3 H_3PO_4
\]

Arsine can be prepared by the reduction of the chloride, (8.3.51) or (8.3.52). The corresponding syntheses can also be used for stibine and bismuthine.

\[
4 AsCl_3 + 3 LiAlH_4 \rightarrow 4 AsH_3 + 3 LiAlCl_4
\]

\[
4 AsCl_3 + 3 NaBH_4 \rightarrow 4 AsH_3 + 3 NaCl + 3 BCl_3
\]

The hydrolysis of calcium phosphide or arsenide can also generate the trihydrides.

**Structure**

The phosphorus in phosphine adopts sp$^3$ hybridization, and thus phosphine has an umbrella structure (Figure 21a) due to the stereochemically active lone pair. The barrier to inversion of the umbrella ($E_a = 155$ kJ/mol) is much higher than that in ammonia ($E_a = 24$ kJ/mol). Putting this difference in context, ammonia’s inversion rate is $10^{11}$ while that of phosphine is $10^3$. As a consequence it is possible to isolate chiral organophosphines (PRR’R”). Arsine adopts the analogous structure (Figure 21b).
Figure 21: The structures of (a) phosphine and (b) arsine.

Reactions

Phosphine is only slightly soluble in water (31.2 mg/100 mL) but it is readily soluble in non-polar solvents. Phosphine acts as neither an acid nor a base in water; however, proton exchange proceeds via the phosphonium ion (PH$_4^+$) in acidic solutions and via PH$_2^-$ at high pH, with equilibrium constants $K_b = 4 \times 10^{-28}$ and $K_a = 41.6 \times 10^{-29}$, respectively.

Arsine has similar solubility in water to that of phosphine (i.e., 70 mg/100 mL), and AsH$_3$ is generally considered non-basic, but it can be protonated by superacids to give isolable salts of AsH$_4^+$. Arsine is readily oxidized in air, (8.3.53).

\[
\text{2 AsH}_3 + 3 \text{O}_2 \rightarrow \text{As}_2\text{O}_3 + 3 \text{H}_2\text{O}
\]

Arsine will react violently in presence of strong oxidizing agents, such as potassium permanganate, sodium hypochlorite or nitric acid. Arsine decomposes to its constituent elements upon heating to 250 - 300 °C.

Gutzeit test

The Gutzeit test is the characteristic test for arsenic and involves the reaction of arsine with Ag$^+$. Arsine is generated by reduction of aqueous arsenic compounds, typically arsenites, with Zn in the presence of H$_2$SO$_4$. The evolved gaseous AsH$_3$ is then exposed to silver nitrate either as powder or as a solution. With solid AgNO$_3$, AsH$_3$ reacts to produce yellow Ag$_4$AsNO$_3$, while with a solution of AgNO$_3$ black Ag$_3$As is formed.

Hazards

Pure phosphine is odorless, but technical grade phosphine has a highly unpleasant odor like garlic or rotting fish, due to the presence of substituted phosphine and diphosphine (P$_2$H$_4$). The presence of P$_2$H$_4$ also causes spontaneous combustion in air. Phosphine is highly toxic; symptoms include pain in the chest, a sensation of coldness, vertigo, shortness of breath, and at higher concentrations lung damage, convulsions and death. The recommended limit (RL) is 0.3 ppm.

Arsine is a colorless odorless gas that is highly toxic by inhalation. Owing to oxidation by air it is possible to smell a slight, garlic-like scent when arsine is present at about 0.5 ppm. Arsine attacks hemoglobin in the red blood cells, causing them to be destroyed by the body. Further damage is caused to the kidney and liver. Exposure to arsine concentrations of 250 ppm is rapidly fatal: concentrations of 25 – 30 ppm are fatal for 30 min exposure, and concentrations of 10 ppm can be fatal at longer exposure times. Symptoms of poisoning appear after exposure to concentrations of 0.5 ppm and the recommended limit (RL) is as low as 0.05 ppm.
Bibliography