If you refer to Table 1 from Ions in Solution (Electrolytes), you will see that pure water does conduct some electrical current, albeit much less than—even the weak electrolytes listed there. This is because water itself is a very weak electrolyte. It ionizes to hydrogen ions and hydroxide ions to an extremely small extent:

\[
\text{H}_2\text{O}(l) \rightleftharpoons \text{H}^+(aq) + \text{OH}^-(aq) \label{1}
\]

Careful measurements show that at 25°C the concentrations of H\(^+\)(aq) and OH\(^-\)(aq) are each 1.005 \(\times\) 10\(^{-7}\) mol dm\(^{-3}\). At higher temperatures more H\(^+\)(aq) and OH\(^-\)(aq) are produced while at lower temperatures less ionization of water occurs. Nevertheless, in pure water the concentration of H\(^+\)(aq) always equals the concentration of OH\(^-\)(aq). Dissolving acids or bases in water can change the concentrations of both H\(^+\)(aq) and OH\(^-\)(aq), causing them to differ from one another. The special case of a solution in which these two concentrations remain equal is called a **neutral solution**.

A hydrogen ion, H\(^+\), is a hydrogen atom which has lost its single electron; that is, a hydrogen ion is just a proton. Because a proton is only about one ten-thousandth as big as an average atom or ion, water dipoles can approach very close to a hydrogen ion in solution. Consequently the proton can exert a very strong attractive force on a lone pair of electrons in a water molecule—strong enough to form a coordinate covalent bond:

\[
\text{H}^+:\ddot{\text{O}}:\text{H} \rightarrow \text{H}:\ddot{\text{O}}:\text{H}^+
\]

The H\(_3\)O\(^+\) formed in this way is called a **hydronium ion** (on the left in the figure below). All three of its O—H bonds are exactly the same, and the ion has a pyramidal structure as predicted by VSEPR theory (1a). To emphasize the fact that a proton cannot exist by itself in aqueous solution, Eq. (1a) is often rewritten as

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

**Figure** (PageIndex{1}) *Hydrogen and hydroxide ions in aqueous solution.* (a) Hydronium ion, H\(_3\)O\(^+\); (b) hydroxide ion, OH\(^-\);

Like other ions in aqueous solution, both hydronium and hydroxide ions are hydrated. Moreover, hydrogen bonds are involved in attracting water molecules to hydronium and hydroxide ions. In both cases three water molecules appear to be rather tightly held, giving formulas H\(_3\)O(H\(_2\)O)\(_3\)^+ (or H\(_9\)O\(_4\)^+) and HO(H\(_2\)O)\(_3\)^- (or H\(_7\)O\(_4\)^-). Possible structures for the hydrated hydronium and hydroxide ions are shown in Fig. (PageIndex{1}).
Hydrogen bonding of hydronium and hydroxide ions to water molecules accounts rather nicely for the unusually large electrical currents observed for some electrolytes containing H and OH. The case of the hydronium ion is illustrated in Fig. \(\PageIndex{2}\). When a hydronium ion collides with one end of a hydrogen-bonded chain of water molecules, a different hydronium ion can be released at the other end. Only a slight movement of six protons and a rearrangement of covalent and hydrogen bonds is needed. In effect a hydronium ion can almost instantaneously “jump” the length of several water molecules. It need not elbow its way through a crowd as other ions must. The same is true of aqueous hydroxide ions. Since both ions move faster, they can transfer more electrical charge per unit time, that is, more current.

**Figure \(\PageIndex{1}\)** (c) hydrated Hydronium ion, \(H_2O_4^+\); (d) hydrated hydroxide ion, \(H_7O_4^-\).

**Figure \(\PageIndex{2}\)** Rapid transfer of protons through aqueous solution. (a) Hydronium ion hydrogen bonded to a chain of water molecules. Minor rearrangement of the positions of H nuclei and hydrogen bonds results in situation (b) where the hydronium ion is at the right end of the chain. (c) An analogous situation would be a row of pickpockets in a crowd. The net result is transfer of a wallet from the person at the left of the row to the right (d). Only the ends of the chain have gained or lost wallets.

**Contributors**

- Ed Vitz (Kutztown University), John W. Moore (UW-Madison), Justin Shorb (Hope College), Xavier Prat-Resina (University of Minnesota Rochester), Tim Wendorff, and Adam Hahn.