The Croonian Lecture

Ionic movements and electrical activity in giant nerve fibres

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[Plate 1]

During the course of an investigation into the central nervous systems of squids and cuttlefish, Professor J. Z. Young (1936a, b) noticed certain transparent tubular structures in the peripheral nerves. These must have seemed too large to be nerve fibres, and in a subsequent article Young (1944) remarked that he first took them to be blood vessels. However, on examining them more closely, he was able to prove that the tubes were, in fact, nerve fibres of exceptional size. Like many important discoveries, this was not an entirely new observation. It had been known since the time of Remak (1843) that Crustacea possessed giant nerve fibres, but with one exception the still larger fibres in cephalopods seem not to have been recognized as such. As Young pointed out, the exception was L. W. Williams who wrote a monograph on the squid which was published in 1909. In this monograph, Williams referred briefly to the large fibres in the nervous system. Williams did not commit himself as to the size of the fibres, but it is clear that he must have seen them. Thus he remarked that

' The very size of the nerve processes has prevented their discovery, since it is well nigh impossible to believe that such a large structure can be a nerve fibre.'

Williams did not follow up this observation and it seems to have passed unnoticed until Young began his investigations in 1933.

In a full grown specimen of the squid, Loligo forbesi, the giant fibres may be as much as 1 mm in diameter and are therefore about fifty times bigger than the largest fibres in man. When isolated from the animal, the fibres continue to conduct electrical impulses which are essentially similar to those found in other nerves. Young realized at once that such a preparation was likely to be of great value to biologists, and it was not long before several physiologists were working with the preparation at Woods Hole in Massachusetts and at Plymouth in England. The main object of this work has naturally been to explain the conduction of nervous impulses in physical terms. At first, electrical methods were the most profitable, but in the years since the war it has been possible to use radioactive tracers to follow ionic movements more directly. The upshot is that we now have a fairly definite picture of the way in which a nerve conducts impulses. This picture is not accepted by everyone and it probably does not apply to all excitable tissues.
Nevertheless, the evidence that most nerves work in essentially the same manner seems sufficiently good to justify the large amount of effort which has been devoted to squid fibres.

**Some general properties of giant nerve fibres**

When viewed under a low-power objective a squid giant fibre appears as a transparent cylinder of protoplasm surrounded by a thin sheath of connective tissue. In the fresh state the protoplasm of cephalopod axons appears to be a gel and when a glass capillary, which has been inserted into the protoplasm, is withdrawn, it leaves a fluid-filled space down which particles can be seen to fall under the influence of gravity.

Figure 1, plate 1, shows an isolated fibre under dark ground illumination. In this case a glass capillary, 100 μ in diameter, has been inserted down the axis of the fibre for a distance of about 30 mm; the capillary can be used either for injecting substances or for recording electrical changes. Provided that the capillary does not touch the surface it does little harm and a large fibre will continue to conduct impulses for many hours after being impaled in this manner.

Another advantage of the giant axon is that samples of protoplasm, or axoplasm as it is called, can be obtained by squeezing out the contents of the axon, in much the same way that one would squeeze out a tube of toothpaste. Since axoplasm obtained in this way is only very slightly contaminated with extracellular fluid, it provides excellent material for chemical analyses. Table 1 summarizes the results of some of these measurements. It will be seen that the axoplasm contains a high concentration of potassium and a low concentration of sodium and chloride. This is the reverse of the situation in the animal’s blood, or in sea water, where sodium and chloride are the dominant ions and potassium is relatively dilute. Potassium ions are probably in the free state inside the fibre and do not seem to be bound to proteins or other large molecules. Thus the mobility and diffusion coefficient of labelled potassium are about the same inside the axon as in free solution (Hodgkin & Keynes 1953), and the membrane potential at which the passive fluxes of potassium are equal, agrees with that calculated from the Nernst equation on the basis that the activity coefficient of potassium ions inside the fibre is approximately equal to that in the external solution (Curtis & Cole 1942; Hodgkin & Keynes 1955b; for evidence in muscle see Adrian 1956; Conway 1957). An interesting point which has come to light recently is that the concentration of calcium in axoplasm is only about 1/30 of that in the external solution (Keynes & Lewis 1956). This figure refers to total calcium and there is evidence from mobility measurements (Hodgkin & Keynes 1957) that most of this calcium must be bound and that the concentration of ionized calcium may be only about 1/1000 of that outside.

The excess of potassium inside the fibre is balanced by organic anions of which, in squid fibres, isethionic acid appears to be the most important (Koelchlin 1955). Other organic compounds which help to balance the cations are aspartic acid and phosphate esters such as adenosine triphosphate, ATP. Isethionic acid is probably not present in vertebrate nerve and here the nature of the anions is still uncertain.
**Figure 1.** Isolated giant axon with 100μ glass capillary inside it, dark-ground illumination; from an experiment of Hodgkin & Keynes (1956).

**Figure 2.** Electronmicrograph of a thin section of two small unmyelinated fibres from the frog’s sciatic nerve, from Robertson (1957). The section was fixed with permanganate. A, axon; S, Schwann cell; M, mitochondrion. Note the membrane at the surface of the axon and Schwann cell, and the space between the axon and Schwann cell. This photograph was kindly provided by Dr Robertson at a time when his paper was in press.

*(Facing p. 2)*
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For our present purpose this may not matter, because there is evidence that sodium and potassium are the ions principally concerned in carrying current through the membrane during the nervous impulse.

At the outside of the fibre there is a thin membrane which acts as a barrier and prevents the ions in the external fluid from mixing rapidly with those in the fibre. This membrane has a high electrical resistance (ca. 1000 ohm cm², Cole & Hodgkin 1939) corresponding to a small but finite permeability to ions and an electrical

**Table 1. Approximate Concentrations of Ions and Other Substances in the Axoplasm of Freshly Isolated Giant Axons and in the External Fluid**

<table>
<thead>
<tr>
<th>Substance</th>
<th>Concentration in Axoplasm</th>
<th>Concentration in Blood</th>
<th>Concentration in Sea Water¹</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>H₂O</td>
<td>865²</td>
<td>870²</td>
<td>966</td>
<td>g/kg</td>
</tr>
<tr>
<td>K</td>
<td>400⁴</td>
<td>20²</td>
<td>10⁵</td>
<td>mmole/kg H₂O</td>
</tr>
<tr>
<td>Na</td>
<td>50⁴</td>
<td>440⁷</td>
<td>460⁶</td>
<td>mmole/kg H₂O</td>
</tr>
<tr>
<td>Cl</td>
<td>40⁸</td>
<td>560⁷</td>
<td>540⁶</td>
<td>mmole/kg H₂O</td>
</tr>
<tr>
<td>Ca</td>
<td>0-4⁹</td>
<td>10⁷</td>
<td>10⁶</td>
<td>mmole/kg H₂O</td>
</tr>
<tr>
<td>Mg</td>
<td>10¹⁰</td>
<td>54⁷</td>
<td>53⁶</td>
<td>mmole/kg H₂O</td>
</tr>
<tr>
<td>Isethionate</td>
<td>270²</td>
<td>—</td>
<td>—</td>
<td>mmole/kg H₂O</td>
</tr>
<tr>
<td>Aspartate</td>
<td>75³</td>
<td>—</td>
<td>—</td>
<td>mmole/kg H₂O</td>
</tr>
<tr>
<td>Glutamate</td>
<td>12³</td>
<td>—</td>
<td>—</td>
<td>mmole/kg H₂O</td>
</tr>
<tr>
<td>Succinate + Fumarate</td>
<td>17²</td>
<td>—</td>
<td>—</td>
<td>mmole/kg H₂O</td>
</tr>
<tr>
<td>Orthophosphate</td>
<td>2-5–9¹¹</td>
<td>—</td>
<td>—</td>
<td>mmole/kg H₂O</td>
</tr>
<tr>
<td>ATP</td>
<td>0-7–1-7¹¹</td>
<td>—</td>
<td>—</td>
<td>mmole/kg H₂O</td>
</tr>
<tr>
<td>Phosphagen</td>
<td>1-8–5-7¹¹</td>
<td>—</td>
<td>—</td>
<td>mmole/kg H₂O</td>
</tr>
</tbody>
</table>

¹ Salinity 3-45 %.
² Koechlin (1955).
³ Robertson (1949).
⁴ Steinbach & Spiegelman (1943); Keynes & Lewis (1951).
⁵ Robertson (1949). See also figure 3, legend.
⁶ Webb (1939, 1940).
⁷ Robertson (1949).
⁸ Steinbach (1941). Value for fresh axoplasm; Koechlin (1955) gives 160 for pooled axoplasm from a large number of squid.
⁹ Keynes & Lewis (1956). Koechlin gives 3-5 but regards this as an upper limit.
¹⁰ Koechlin (1955). This is regarded as an upper limit.
¹¹ Caldwell (1956).

Concentrations in axoplasm have been calculated from the original authors' figures on the basis that axoplasm contains 865 g H₂O per kg axoplasm. For similar data in other fibres see Hodgkin (1951).

capacity of about 1 μF/cm² (Curtis & Cole 1938). These are the values which might be expected from a bimolecular layer of lipid with a thickness of 50 Å, a dielectric constant of about 5 and an electrical resistivity of 2 x 10⁹ ohm cm. The high resistivity of the membrane is in sharp contrast to the resistivities of the axoplasm and external fluid, which are about 30 and 20 ohm cm, respectively (Cole & Hodgkin 1939). Under the electron-microscope the surface membrane of fibres fixed in osmic acid or permanganate appears as a layer 50 to 100 Å thick (Geren & Schmitt 1954; Robertson 1957). Figure 2, plate 1, is an electronmicrograph obtained recently by Dr Robertson; it shows the thin membrane at the surface of
two small unmyelinated fibres from the frog, and the Schwann cells which surround but do not completely enclose the fibres. In the squid axon, the structure of the Schwann cell layer is more complicated, but indirect evidence suggests that this layer may be only a relatively weak barrier to ionic movement, and that the main restriction occurs at the 50 to 100 Å membrane at the surface of the axon (see Frankenhaeuser & Hodgkin 1956).

<table>
<thead>
<tr>
<th>animal</th>
<th>axon diameter (µm)</th>
<th>temperature (°C)</th>
<th>Na entry</th>
<th>K loss</th>
<th>method</th>
<th>references</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Carcinus maenas</em></td>
<td>30</td>
<td>ca. 17</td>
<td>—</td>
<td>1·7</td>
<td>indirect</td>
<td>Hodgkin &amp; Huxley (1947)</td>
</tr>
<tr>
<td><em>C. maenas</em></td>
<td>30</td>
<td>ca. 17</td>
<td>—</td>
<td>2·4</td>
<td>indirect 42K</td>
<td>Keynes (1951a)</td>
</tr>
<tr>
<td><em>Sepia officinalis</em></td>
<td>200</td>
<td>ca. 15</td>
<td>—</td>
<td>3·4</td>
<td>indirect</td>
<td>Weidmann (1951)</td>
</tr>
<tr>
<td>S. officinalis</td>
<td>200</td>
<td>ca. 15</td>
<td>3·7*</td>
<td>4·3†</td>
<td>activation analysis 42K and 24Na</td>
<td>Keynes (1949, 1951b)</td>
</tr>
<tr>
<td>L. pealii</td>
<td>500</td>
<td>ca. 22</td>
<td>3·8</td>
<td>3·6</td>
<td>activation analysis</td>
<td>Keynes &amp; Lewis (1951)</td>
</tr>
<tr>
<td>L. pealii</td>
<td>500</td>
<td>ca. 22</td>
<td>5·5</td>
<td>3·0</td>
<td>activation analysis</td>
<td>Keynes &amp; Lewis (1951)</td>
</tr>
<tr>
<td>L. pealii</td>
<td>500</td>
<td>room temperature</td>
<td>4·6</td>
<td>—</td>
<td>24Na</td>
<td>Rothenberg (1950)</td>
</tr>
<tr>
<td>L. pealii</td>
<td>500</td>
<td>24</td>
<td>—</td>
<td>3·1</td>
<td>flame photometry</td>
<td>Grundfest &amp; Nach-mansohn (1950)</td>
</tr>
<tr>
<td>L. pealii</td>
<td>500</td>
<td>6</td>
<td>—</td>
<td>9·3</td>
<td>flame photometry</td>
<td>Shanes (1954)</td>
</tr>
</tbody>
</table>

* Activity increases both the influx and the efflux of Na (as is to be expected if the membrane potential approaches the equilibrium potential for Na); the figure of 3·7 is the difference between the extra influx of 10·3 and the extra efflux of 6·6 pmole/cm² per impulse.
† Corresponding figures for K are extra influx 0·4, extra efflux 4·7 pmole/cm² per impulse.

During an impulse the conductance of the surface membrane increases about 40-fold (Cole & Curtis 1939) and sodium and potassium ions cross the membrane more easily than in the resting state (Keynes 1951b). The net effect is that each impulse is associated with a small inward movement of sodium ions and a small outward movement of potassium ions. These movements have been measured in various ways and the different methods agree in giving the quantity which crosses one square centimetre of membrane in one impulse as 3·4 × 10⁻¹² mole (20° C, cephalopod axons; see table 2). This is about 20000 ions per square micron of surface.

From an electrical point of view the nervous impulse consists of a brief change in membrane potential which travels down the fibre at a velocity of about 20 m/sec. Figure 3, which was contained in collaboration with Dr Keynes, illustrates the form of the action potential in intact and isolated axons from the squid. The record on the left was made by inserting a microelectrode with a tip diameter of about 0·5 µ into an intact axon in a squid which had been subjected to only slight operative procedure; the natural circulation was still intact. When the microelectrode was pushed through the surface of the axon, the potential jumped suddenly to a new value about 70 mV negative to the zero. This illustrates the well-known point that there is a resting potential and that the inside of a nerve fibre is normally negative
to the outside. When the axon was stimulated with an electric shock, an impulse travelled along it; the inside swung momentarily positive and then returned to its resting level. The total amplitude of the action potential was about 110 mV and its duration (at 8.5°C) about 1.2 msec. In this case since we were working with the intact animal, the nerve was still connected to the muscles. A few milliseconds after the impulse had travelled past the microelectrode it reached the muscles; the body wall then gave a powerful flap which smashed the microelectrode and almost certainly damaged the axon. For this and many other reasons it is simpler to work with isolated axons. The action potential of an isolated fibre is shown in figure 3B; it differs slightly from the one seen in the intact animal, but the essential features of the conduction process appear to be the same in both cases.

![Graph showing action potentials](image)

**Figure 3.** Action potentials in intact (A) and isolated axon (B) from Loligo forbesi. The records are taken from an unpublished series of experiments carried out by Dr R. D. Keynes and the author in 1952.

A fresh and lively squid was taken out of the aquarium and immobilized by cutting the nerves connecting the head with the stellate ganglion. The mantle was opened ventrally by a single cut and was spread out under cooled oxygenated sea water in a transparent dish. Using a motor car headlamp to illuminate the animal and a binocular microscope to watch the penetration, a microelectrode with a tip diameter of about 0.5 μ was pushed through the muscle and into the hindmost giant nerve fibre. Only small and transient changes in potential occurred as the electrode passed through the muscle but a step of 70 mV occurred when it penetrated the giant fibre; no twitch occurred at this stage. The action potential was elicited by applying a shock to the central part of the giant axon. Record B was obtained later (with a similar electrode) in the corresponding axon from the other side of the same squid; this axon had been isolated in the usual way and was immersed in sea water.

The following values were obtained in five successful experiments of this type: resting potential, 63 to 72 mV, mean 68 mV; action potential, 99 to 115 mV, mean 107 mV; hyperpolarization after spike, 1 to 7 mV, mean 4 mV.

The potassium concentrations in five samples of blood from other squid which had been immobilized in a similar way was 17.7 to 25.8 mmole/kg H₂O, mean 19.6 mmole/kg H₂O.

Rather more than 50 years ago, Bernstein (1902) suggested that the resting membrane was selectively permeable to potassium ions and that the potential difference across it arose from the tendency of potassium ions to move outwards from the more concentrated solution inside a nerve or muscle fibre. Thus, if a membrane which is selectively permeable to potassium separates axoplasm containing 400 mM-K from plasma containing 20 mM-K, one would expect the inside
of the membrane to be 75 mV negative to the external solution. This value is
obtained from the Nernst relation
$$V_K = \frac{RT}{F} \ln \frac{[K]_o}{[K]_i},$$
(1)
where $V_K$ is the equilibrium potential of the potassium ion defined in the sense, internal potential minus external potential, $[K]_o$ and $[K]_i$ are the potassium concentrations (or strictly activities) outside and inside the fibre and $R$, $T$ and $F$ have their usual significance. In an undissected squid axon, with natural circulation, resting potentials of about 70 mV have been observed by Moore & Cole (1955) and by the author and Dr Keynes (figure 3). In muscle, where equation (1) predicts a value of 102 mV for the potassium equilibrium potential, the resting potential is 90 to 95 mV (Adrian 1956).

The resting potential of an isolated squid axon in sea water containing 10 mm-K is only 50 to 60 mV, whereas the potassium equilibrium potential should be about 90 mV. The reason for the low resting potential in isolated axons is not fully understood but possible factors may be (1) that small branches have to be cut through in isolating the axon, (2) the ends of the fibre are cut, (3) the fluid used to bathe the isolated fibres is different from that in which they are naturally immersed. The large difference between the potassium equilibrium potential and the resting potential in isolated fibres probably accounts for the large underswing, or transient phase of hyperpolarization, which follows the main part of the spike in isolated fibres. During the later stages of the spike, the potassium permeability rises and the state of increased potassium permeability persists for several milliseconds after the spike. The effect of this is to make the membrane potential approach more closely to the potassium equilibrium potential, with the result that the fibre undergoes a transient phase of hyperpolarization. Evidence for this suggestion is provided by the observation that the membrane potential during the underswing is affected by potassium concentrations which have relatively little effect on the resting membrane potential (Hodgkin & Katz 1949; Hodgkin & Keynes 1955; Frankenhaeuser & Hodgkin 1956). In other words, during the underswing equation (1) holds down to lower potassium concentrations than it does in the resting state (Hodgkin & Keynes 1955).

In support of Bernstein's idea it is found that the resting potential disappears if the external potassium concentration is made equal to the internal concentration and that, except at low external concentrations of potassium, the potential varies in the manner predicted by equation (1) (see Curtis & Cole 1942; Adrian 1956; Hodgkin & Keynes 1955). However, these experiments also show that at low concentrations potassium is not the only ion which carries current through the membrane, for the variation with external potassium concentration is less steep in the physiological region than one would expect from (1). At present we do not know what other ions besides potassium contribute to the resting potential in nerve. Since chloride ions are more concentrated outside the fibre they would tend to make the inside of the fibre negative to the outside. In muscle, where the distribution of chloride and potassium ions closely resembles that in a Donnan system (Boyle & Conway 1941), there is evidence that both ions contribute to the potential and
that, for very small displacements of potential, the transport numbers are approximately 0.3 for K and 0.6 for Cl (unpublished experiments with Dr P. Horowicz).

During the nervous impulse the internal potential swings momentarily from its resting value of $-70 \text{ mV}$ to one of about $+40 \text{ mV}$ (figure 3.4). Activity therefore involves a reversal of potential difference and so cannot depend on a breakdown of the membrane as Bernstein supposed. A simple way of explaining the reversal of membrane potential is to assume that when the membrane is activated by an electric current it momentarily becomes selectively permeable to sodium ions. On this basis the potential difference at the crest of activity arises from the tendency of the sodium ions to move into the fibre from the more concentrated solution outside. In support of this idea it is found that the action potential of many but not all excitable tissues disappears in the absence of external sodium or lithium ions, and that the reversed potential difference across the active membrane varies with external sodium concentration in the same manner as a sodium electrode. The state of increased sodium permeability wears off after about a millisecond and the potential therefore tends to return to its original level. The process of repolarization is greatly accelerated by the fact that depolarization causes a delayed rise in the permeability to potassium ions. This speeds up the exit of potassium and keeps the whole action potential reasonably short.

The effect of sodium concentration on the action potential

The importance of sodium in maintaining excitability was first recognized by Overton (1902) who suggested that impulse conduction was accompanied by an exchange between intracellular potassium ions and extracellular sodium ions. References to papers dealing with the effect of sodium on the action potentials of different excitable tissues are given in table 3.

Deviations from the typical sodium electrode type of behaviour have been seen in turtle ventricle, and in frog muscles treated with tetraethylammonium ions. In both cases the fibres fail in the complete absence of sodium, but at intermediate concentrations the overshoot changes much less than would be expected from the simple theory. Crab muscle fibres, previously treated with tetrabutylammonium ions, give a large overshoot in the absence of sodium or any other monovalent cation so that the sodium theory clearly does not apply in this case. However, in the majority of excitable fibres, the effects of sodium-deficient solutions agree well with the hypothesis that the action potential depends on an increase in sodium permeability.

The relation between sodium concentration and overshoot has been investigated with solutions containing extra sodium chloride as well as those in which choline replaces sodium, or sucrose replaces sodium chloride. Since sodium-rich solutions increase the spike, it cannot be argued that their action is due to some kind of reversible damage or narcosis of the membrane. An extreme example of the action of added NaCl is afforded by an experiment mentioned by Stämpfli (1956) in which a sudden four-fold increase of NaCl raised the overshoot of a node of Ranvier by about 38 mV. In carrying out experiments with solutions containing extra sodium it is important either to measure the action potential very soon after changing the
solution, or to make the comparison between isosmotic solutions, for example, between one solution containing extra sodium chloride and another containing an osmotically equivalent amount of sucrose or choline chloride. If this is not done, the effect of the high external sodium will be neutralized by the rise in internal sodium concentration which results from the removal of water by the hypertonic

Table 3. Effect of sodium ions on the action potentials of different tissues—summary and references

<table>
<thead>
<tr>
<th>preparation</th>
<th>substance used to replace Na or NaCl</th>
<th>rapid reversible block in Na-free solution</th>
<th>active membrane behaves like a sodium electrode, i.e. slope of 58 mV ± ca. 5 mV for 10-fold change</th>
<th>references</th>
</tr>
</thead>
<tbody>
<tr>
<td>squid giant axon</td>
<td>choline, dextrose</td>
<td>yes</td>
<td>yes</td>
<td>Hodgkin &amp; Katz (1949), Cole (1955)</td>
</tr>
<tr>
<td>Sepia giant axon</td>
<td>choline</td>
<td>yes</td>
<td>—</td>
<td>unpublished</td>
</tr>
<tr>
<td>crab (Carcinus) nerve fibre</td>
<td>choline, dextrose, sucrose</td>
<td>yes</td>
<td>—</td>
<td>Katz (1947), Hodgkin &amp; Katz (1949)</td>
</tr>
<tr>
<td>crab muscle</td>
<td>choline, and other quaternary ammonium salts choline*, dextrose</td>
<td>no</td>
<td>no</td>
<td>Fatt &amp; Katz (1953)</td>
</tr>
<tr>
<td>frog single myelinated nerve fibre*</td>
<td>choline, sucrose</td>
<td>yes</td>
<td>yes</td>
<td>Huxley &amp; Stämpfli (1949, 1951)</td>
</tr>
<tr>
<td>frog sartorius muscle</td>
<td>choline, sucrose</td>
<td>yes</td>
<td>yes</td>
<td>Overton (1902), Nastuk &amp; Hodgkin (1950), Hagiwara &amp; Watanabe (1955)</td>
</tr>
<tr>
<td>frog sartorius muscle</td>
<td>tetraethylammonium</td>
<td>yes</td>
<td>no</td>
<td>Hagiwara &amp; Watanabe (1955)</td>
</tr>
<tr>
<td>dog Purkinje fibres</td>
<td>sucrose</td>
<td>yes</td>
<td>yes</td>
<td>Draper &amp; Weidmann (1951)</td>
</tr>
<tr>
<td>frog ventricle</td>
<td>choline, sucrose</td>
<td>yes</td>
<td>yes</td>
<td>Brady &amp; Woodbury (1957)</td>
</tr>
<tr>
<td>turtle ventricle</td>
<td>choline after atropine</td>
<td>yes</td>
<td>no</td>
<td>S. Weidmann (unpublished)</td>
</tr>
<tr>
<td>guinea-pig ventricle</td>
<td>choline</td>
<td>yes</td>
<td>no</td>
<td>Coraboeuf &amp; Otsuka (1956)</td>
</tr>
</tbody>
</table>

For the action of guanidine and other quarternary ammonium compounds in frog nerve see Larramendi, Lorente de Nó & Vidal (1956), Lorente de Nó, Vidal & Larramendi (1957) and Lorente de Nó (1949).

* Müller (1956) reports prolonged action potentials in Na-free choline in a node which had previously been depolarized with a large outward current.

solution. This point seems to have been overlooked by Shaw, Simon, Johnstone & Holman (1956) and may help to account for the difference between their conclusions and those of other workers. Thus Nastuk & Hodgkin (1950, table 4) found overshoots of 33, 25 and 33 mV in frog muscle fibres immersed in Ringer’s fluid, Ringer plus 100 mm-sucrose and Ringer plus 50 mm-NaCl, respectively. Shaw et al. (1956) found about the same overshoot in Ringer’s fluid or Ringer’s fluid plus 26 to 60 mm-NaCl, but they apparently did not carry out any control experiments with Ringer’s fluid plus sucrose or choline chloride.

Evidence about the relation between the internal sodium concentration and overshoot is somewhat conflicting. Desmedt (1953) found that the active membrane behaved like a sodium electrode except when the internal sodium concen-
tration was low, in which case the contribution of other ions might no longer be small compared with that of the sodium ion. On the other hand Shaw, Simon & Johnstone (1956) and Shaw et al. (1956) consider that there is no correlation between the overshoot of toad muscle and the internal sodium concentration calculated on the assumption that the extracellular space occupies 15% of the muscle volume. In the writer’s opinion the difference between the two conclusions may result from the very great difficulty of making accurate measurements of internal sodium concentration in a tissue in which much of the total sodium may be in the extracellular space. In the squid giant axon, raising the internal sodium by micro-injection reduced the overshoot by approximately the amount expected from the Nernst equation (Hodgkin & Keynes 1956).

**The local circuit theory**

While there is still much argument about the nature of the action potential, everyone is agreed that conduction of the nervous impulse is essentially an electrical process and that propagation is brought about by current flowing in a local circuit at the front of the action potential. In explaining how the impulse propagates, it

![Diagram](image_url)

**Figure 4.** Diagrams illustrating the local circuit theory; the upper sketch represents an unmyelinated nerve fibre, the lower a myelinated nerve fibre. For evidence about saltatory conduction in myelinated fibres see numerous papers by Tasaki and colleagues (reviewed in Tasaki 1953); Huxley & Stämpfli (1949); Stämpfli (1952).

is convenient to adopt the view outlined in the first part of this article, and to assume that when the membrane potential is reduced beyond a certain level the membrane changes from a stage of moderate potassium permeability to one of high sodium permeability. This assumption is made for the sake of definiteness; it is not essential to the local circuit theory, which can be formulated without specifying how the action potential arises.

The upper part of figure 4 illustrates the flow of current in an unmyelinated axon, of which the squid giant fibre is an example. Suppose that point $A$ is active and that point $B$ is resting. $A$ is sodium permeable so the inside of the fibre is positive; $B$ is potassium permeable so the inside is negative. Current therefore flows in a local circuit between resting and active nerve. It will be seen that current flows in the same direction as the impulse in the axis cylinder and returns in the opposite direction in the external fluid. This current reduces the membrane potential just ahead of the active region by drawing charge out of the membrane capacity. When
the potential difference has been reduced by about 20 mV the permeability to sodium rises and the inside of the fibre becomes positive. Point B is now active and can stimulate the next region in precisely the same manner. In this way, a wave of increased sodium permeability propagates along the fibre.

This is a convenient point at which to draw attention to an important difference between our own nerves and those of the squid. In most invertebrates, the nerve fibres are continuous unmyelinated structures and the impulse spreads smoothly from one point to the next. Owing to the large capacity of the surface membrane and the resistance of the axon cylinder, this type of conduction only gives a high velocity if the fibre is large. In the myelinated fibres, which form the bulk of our own nerves, a neater method of getting a high velocity has been evolved. These fibres are coated over most of their length with a layer of myelin which acts as an insulator. The excitable membrane is exposed only at certain points known as nodes of Ranvier. When one node becomes active, current flows in a local circuit through the next node in the manner shown in figure 4. Some current is also wasted in charging up the myelin, but since the myelin sheath is relatively thick its capacity is much less than that of an excitable membrane. The effect of this type of conduction, which is known as saltatory conduction, is that in myelinated fibres the impulse is conducted both at greater velocity and with greater economy than in unmyelinated fibres of comparable size.

**Current-voltage relations of the membrane; the voltage-clamp method**

The electrical properties of the membrane have been studied quantitatively by a technique, known as the voltage-clamp method, in which the membrane potential is suddenly displaced from its resting value and held at a fixed potential by a feedback amplifier. The current which flows through a definite area of membrane under the influence of the impressed voltage is measured with a separate amplifier (see Cole 1949; Hodgkin, Huxley & Katz 1952).

In the experiment of figure 5, the internal potential of the fibre was raised suddenly by 65 mV by applying a rectangular pulse to the feedback amplifier which controlled the membrane potential. Since the inside of the fibre was normally at about −50 mV the effect of the pulse was to swing the membrane potential from −50 to +15 mV. The lower records show the current which flowed through the membrane in the presence and absence of external sodium ions.

When the inside of the fibre is made more positive the potential difference and charge on the membrane is reduced, or reversed if the pulse is large enough. The immediate effect of the sudden change is that a large surge of capacity current flows through the membrane. This surge is complete in about 20 μsec and is too rapid to show on records such as those in figure 5. The surge is proportional to the

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* Convention as to signs. In this article membrane potentials are given in the sense internal potential minus external potential; outward currents are therefore taken as positive and are shown as upward deflexions. Although this convention seems the more logical and is widely used, I have previously hesitated to adopt it because it conflicts with the accepted use of words such as negative after-potential. However, since many workers now use internal electrodes and since the action potential is almost invariably shown as an upward deflexion, it seems right to make a change.
amplitude of the step and is little influenced by changes in temperature or ionic concentration. Quantitatively, it corresponds to the charge or discharge of a condenser with a capacity of 0.9 μF/cm² (Hodgkin et al. 1952); this value is in reasonable agreement with that obtained by other methods (Curtis & Cole 1938).

The direction and amplitude of the 'ionic current' which flows through the membrane during the period of maintained depolarization varies greatly with the amplitude of the voltage step and with the concentration of ions in the external medium. In sea water, which contains about 460 mm-Na, the initial phase of ionic current is inward for depolarizations of 15 to 100 mV in spite of the fact that such pulses make the internal potential rise. This component of the current is therefore in the opposite direction to that expected in a stable system. If the inward current had not been drawn off by the feedback amplifier it would have continued to make the inside of the fibre more positive at about the same rate as in an action potential. A simple explanation of the inward or reverse current is that when the membrane is depolarized, it undergoes a rise in sodium permeability, which allows sodium ions to move inwards from the more concentrated solution outside. In support of this idea it is found that if an inert cation, such as choline, is substituted for sodium in the external medium, the inward current disappears and is replaced by a corresponding hump of outward current (figure 5). Under

**Figure 5.** Membrane currents associated with depolarization of 65 mV in presence and absence of external sodium ions. The change in membrane potential is shown at the top; the lower three records give the membrane current density; temperature, 11° C; outward current and internal potential shown upward. From Hodgkin & Huxley 1952a; note that the records have been reproduced so as to show outward current upwards and not downwards as in the original article.
these conditions there are no sodium ions in the external solution and the outward current arises from the tendency of the sodium ions in the axoplasm to move to the sodium-free solution outside.

If the sodium concentration is kept at its normal value and the amplitude of the voltage step is increased to, say, 140 mV the initial current again becomes outwardly directed (figure 6). The reason, presumably, is that the inside of the fibre is now sufficiently positive to overcome the tendency of the sodium ions to move inwards from the more concentrated solution outside. When the potential step is

\[ V_{Na} = \frac{RT}{F} \ln \frac{[Na]^o}{[Na]_t}, \]  

Figure 6. Membrane currents associated with large depolarizations; axon in sea water, temperature 3-5°C; outward current upwards. The displacement of the membrane potential from its resting value is shown at the right (from Hodgkin et al. 1952).

Figure 7. Diagram of an element of the excitable membrane of a nerve fibre: \(a\), constant capacity; \(b\), channel for Na; \(c\), channel for K; \(d\), channel for other ions (e.g. Cl). After Huxley (1954).

given a certain value, 117 mV in the experiment of figure 6, the initial hump of sodium current disappears and the current then remains at a small and practically constant value for a period of about 300 \(\mu\)sec at 3-5°C. The implication is that there is a definite and fixed equilibrium potential at which one component of the membrane current is zero. Thus, in the scheme shown in figure 7, decreases in the resistance \(R_{Na}\) will produce outward currents for \(V>V_{Na}\), inward currents for \(V<V_{Na}\) and no change in current for \(V=V_{Na}\). If the early hump of current is carried by sodium ions, the equilibrium potential, at which there is no sodium current, should be given by

\[ V_{Na} = \frac{RT}{F} \ln \frac{[Na]^o}{[Na]_t}, \]
where $V_{Na}$ is the equilibrium potential of the sodium ion, defined as an internal potential, $[Na]_o$ and $[Na]_i$ are the sodium concentrations (or strictly activities) outside and inside the fibre. The validity of (2) was tested by measuring the relation between $[Na]_o$ and $V_{Na}$ using choline as a substitute for sodium; the results (Hodgkin & Huxley 1952a, table 1) showed that the equation holds to within about 1 mV over a ten-fold range of $[Na]_o$.

After a short time, about 500 $\mu$sec in the experiment of figure 6, the initial current (which we may now call sodium current) declines, and is replaced by a second component which is always outwardly directed for voltage steps in which the inside of the fibre becomes more positive. The second component can be seen uncomplicated by sodium current if the membrane potential during the voltage step is made equal to $V_{Na}$ (117 mV below the resting potential in the experiment of figure 6). Here the current changes little at first and then increases along an S-shaped curve to an approximately steady value of about 3 mA/cm$^2$. This component of the current is not much altered if sodium ions are replaced by choline and is also present in chloride-deficient solutions. Since potassium ions are known to move outwards during the spike it seemed likely that the second, or delayed, component of the current, was carried by potassium ions. In support of this idea, subsequent experiments with tracers showed that depolarization caused a large increase in the efflux of potassium from Sepia axons and that the quantity of potassium ions crossing a given area of membrane in a given time was equivalent to the total electric charge passing through the same area of membrane in the same time (Hodgkin & Huxley 1953; see also figure 8 of this article). Further evidence is that the equilibrium potential of the second component varies with the external potassium concentration, though not as steeply as predicted by the Nernst equation. Later work (Frankenhaeuser & Hodgkin 1956) suggests that the discrepancy is at least partly explained by the fact that potassium ions tend to accumulate in a space outside the membrane, and that the effective concentration immediately outside the potassium-sensitive layer may not be the same as that in the external solution.

Having established that the early current was probably carried by sodium ions, and the delayed current by potassium ions, the next step was to separate the total current into its two components. The method is illustrated by figure 9. Here curve A shows the ionic current when an axon immersed in sea water was depolarized by 56 mV; under these conditions both sodium and potassium ions contribute to the current. In principle, sodium current can be eliminated by reducing the sodium concentration until the sodium potential is at 56 mV. In practice it would be difficult to hit off exactly the right sodium concentration so the procedure was to interpolate between curves in high and low sodium. Curve B in figure 9, which was in fact very close to the record in 46 mm-Na, was obtained in this way and is taken as the potassium current. If it is assumed (and this may not be quite correct) that the potassium current is independent of the sodium and choline concentrations, then the sodium current in record A can be obtained by subtracting curve B from curve A. This method works well if the sodium current is reasonably large compared to the potassium current, but, as might be expected, it gives errors and anomalies when the former is small compared with the latter.
A convenient way of expressing the permeability of the membrane is to calculate sodium or potassium conductances, \( g_{Na} \) or \( g_{K} \), by means of the relations

\[
g_{Na} = \frac{I_{Na}}{V - V_{Na}},
\]

or

\[
g_{K} = \frac{I_{K}}{V - V_{K}}.
\]

![Graph](image)

**Figure 8.** Relation between membrane current density and potassium efflux when a *Sepia* axon is depolarized (from Hodgkin & Huxley 1953). The axon was depolarized by an applied current for periods of 60 to 600 sec. Vertical lines show \( \pm 2 \times \text{s.e.} \); the horizontal line is drawn at a level corresponding to complete suppression of the average resting efflux.

Subsequent experiments (Hodgkin & Keynes 1955b) indicate that the influx of K remains at a low value when the fibre is depolarized by applied currents; hence the increase in efflux shown above should be very nearly equal to the total outward movement of K.

These definitions would be valid whatever the relations between \( I_{Na} \) and \( V - V_{Na} \), or between \( I_{K} \) and \( V - V_{K} \). However, the usefulness of the definitions is increased by the observation that for sudden changes in potential, the instantaneous value of the current, \( I_{Na} \) or \( I_{K} \), is directly proportional to the driving force, \( V - V_{Na} \) or \( V - V_{K} \).

The continuous curves in figure 10 illustrate the way in which the conductances change when the fibre is depolarized by 56 mV. The sodium conductance rises rapidly to a peak and then declines, in spite of the fact that the fibre is held in the
**Figure 9.** Separation of current into components carried by Na and K, from Hodgkin & Huxley (1952a, figure 5). A depolarization of 56 mV was applied at t=0; the temperature was 8.5°C. Outward current is shown upwards.

---

**Figure 10.** Changes in sodium and potassium conductance associated with depolarization of 56 mV, applied at t=0; temperature 8.5°C. The continuous curves, which were obtained from the experiment of figure 9, show the changes in conductance when the depolarization was maintained; the broken curves, which were drawn from data obtained in other experiments, give the effect of repolarizing the fibre after 0.6 or 6.3 msec. From Hodgkin & Huxley (1952a, b, d).
depolarized state. The potassium conductance changes little at first and then rises, along an S-shaped curve to a level which is maintained as long as the fibre is kept depolarized. The broken lines in figure 10 give the effect of repolarizing the membrane at different times and show that the changes in conductance are reversible. In contrast to the S-shaped rise in conductance associated with depolarization, both sodium and potassium conductances return to their resting values along

![Graph showing changes in sodium and potassium conductance](image)

**Figure 11.** Changes in sodium and potassium conductance associated with different depolarizations at 6°C. The numbers attached to the curves give the depolarizations used. The circles are experimental estimates and the smooth curves are solutions of the equations used to describe the changes in conductance. In a similar figure, reproduced in Hodgkin & Huxley (1952e, f) the potassium curves were drawn 25% too large on the vertical scale. The results were plotted correctly in the original article (Hodgkin & Huxley 1952d) and the error has been corrected in the figure shown above.

approximately exponential curves when the membrane is repolarized. For sodium, the time constant is 0.1 to 0.2 msec at 6°C; for potassium it is 5 to 10 msec. The inactivation process which leads to a decline in sodium conductance during a maintained depolarization is also reversed by repolarizing the membrane and has a time constant of about 10 msec at 6°C. The rate constants governing the turning-on and turning-off of the sodium or potassium conductance have been shown to depend on temperature, membrane potential and calcium concentration, but are little affected by changes in sodium concentration, or by the direction of the ionic current (Hodgkin & Huxley 1952a−d; Frankenhaeuser & Hodgkin 1957, for the effect of Ca).
The changes in sodium and potassium conductances at various depolarizations are given in figure 11. The amplitude and time course of the two conductances vary greatly with membrane potential, but there is no sudden break or discontinuity in the relation between conductance and potential.

Quantitative description of the changes in membrane conductance

The results described in the preceding paragraphs suggest that reducing the membrane potential leads to a transient increase in sodium conductance and a delayed increase in potassium conductance. At first sight these changes appear too simple to account for the varied and complicated reactions of a nerve fibre to electrical stimuli. In order to examine this point, Huxley and I developed a series of equations which described the changes in conductance with reasonable accuracy (Hodgkin & Huxley 1952d). The main object of the equations was to obtain an empirical description of the way in which the permeability depends on time and membrane potential. However, it is perhaps worth mentioning the general type of physical picture which we had in mind in formulating the equations.

To account for the change in potassium conductance we assumed that a path for potassium was formed when four charged particles had moved to a certain region of the membrane under the influence of the electric field. If \( n \) is the probability that a single particle is in the right place, then \( g_K = \bar{g}_K n^4 \), where \( \bar{g}_K \) is the maximum potassium conductance. The value of \( n \) is given by

\[
\frac{dn}{dt} = \alpha_n (1 - n) - \beta_n n,
\]

where \( \alpha_n \) and \( \beta_n \) are rate constants which, at a fixed temperature and calcium concentration, depend only on the membrane potential, \( V \). \( \alpha_n \) increases and \( \beta_n \) decreases as the inside of the fibre becomes more positive.

For the sodium channel we assumed that three simultaneous events each of probability \( m \), opened the channel to Na and that a single event of probability \( (1 - h) \) blocked it. These events need not be specified, but may be thought of as the movement of three activating particles and of one blocking particle to a certain region of the membrane. The probability that there will be three activating particles and no blocking particle is therefore \( m^3 h \). Hence \( g_{Na} = \bar{g}_{Na} m^3 h \), where \( \bar{g}_{Na} \) is the maximum sodium conductance. The values of \( m \) and \( h \) are given by relations similar to (3), i.e.

\[
\frac{dm}{dt} = \alpha_m (1 - m) - \beta_m m,
\]

\[
\frac{dh}{dt} = \alpha_h (1 - h) - \beta_h h.
\]

The \( \alpha \)'s and \( \beta \)'s in these equations depend on temperature, calcium concentration and membrane potential. The effect of making the inside of the fibre more positive is to increase \( \alpha_m \) and \( \beta_h \) and to decrease \( \beta_m \) and \( \alpha_h \).

It is relatively simple to apply equations (3), (4) and (5) to the voltage-clamp data. At a fixed voltage the \( \alpha \)'s and \( \beta \)'s are constant so equations (3), (4) and (5)
lead to exponential expressions for \( n, m \) and \( h \); conductances are then calculated from the relations \( g_k = \bar{g}_k n^4 \) and \( g_{Na} = \bar{g}_{Na} m^3 h \). With values of \( \alpha \) and \( \beta \) which change with membrane potential in a consistent manner, a fair though not a perfect fit is obtained. This is illustrated by figure 11 in which the smooth curves have been calculated and the circles are experimental estimates of the sodium and potassium conductance. The effects of repolarizing the membrane in shutting-off the conductances were taken into account in formulating the equations, and are described by them without any further assumption.

The complete expression for the membrane current density, \( I \), is

\[
I = c \frac{\partial V}{\partial t} + (V - V_K) \bar{g}_K n^4 + (V - V_{Na}) \bar{g}_{Na} m^3 h + (V - V_L) g_L. \tag{6}
\]

The first term on the right-hand side is the capacity current, \( c \) being the membrane capacity per unit area. The second and third terms give the potassium and sodium currents while the last term, which is relatively unimportant, gives the current carried by ions other than sodium and potassium through a leak conductance \( \bar{g}_L \).

If an action potential is elicited over a length of nerve, by applying a short shock to a long metal wire in the axoplasm, there is no longitudinal current and no radial current at any time after the shock. Under these conditions (6) can be simplified by putting \( I = 0 \) for \( t > 0 \), the boundary condition being the initial displacement of \( V \). Solutions of this kind have been worked out numerically, or with an automatic computer; the results agree reasonably with the behaviour of a real nerve. The case in which \( I = \) constant has been calculated by Cole (1954) and shows that, above a certain strength of current, the theoretical membrane gives a train of spikes, which again are not unlike those in a real nerve.

To calculate the form and velocity of the propagated action potential equations (3) to (6) are used in conjunction with the well-known relation for the current density in a continuous nerve fibre surrounded by a large volume of external fluid, that is

\[
I = \frac{a}{2R} \frac{\partial^2 V}{\partial x^2}, \tag{7}
\]

where \( a \) is the radius of the axoplasm and \( R \) is its resistivity; \( x \) is distance along the nerve. In the case of a fibre propagating at constant velocity (\( \theta \)), \( x \) may be replaced by \(-\theta t\).

Hence

\[
\frac{a}{2R\theta^2} \frac{d^2 V}{dt^2} = c \frac{dV}{dt} + (V - V_K) \bar{g}_K n^4 + (V - V_{Na}) \bar{g}_{Na} m^3 h + (V - V_L) g_L. \tag{8}\]

In this equation, the conduction velocity, \( \theta \), is constant, but its value is unknown at the beginning of the computation. The procedure is to guess a value for \( \theta \) and start a trial solution. It is found that \( V \) goes to \( \pm \infty \) according to whether \( \theta \) has been chosen too high or too low. The correct value of \( \theta \), which corresponds to the natural velocity of propagation, brings the potential back to its resting value at the end of the run. A numerical solution along these lines was worked out by Mr Huxley and was found to agree reasonably with the behaviour of a real nerve.
in the following respects: (1) the form, amplitude and velocity of the action potential, (2) the time course and amplitude of the conductance change, (3) the total movements of Na and K during the spike. Huxley's solution which has since been checked by Cole and his colleagues (personal communication) using an automatic computer is shown in figure 12. The conclusion from the analysis is that reversible changes in sodium and potassium conductance, of the kind deduced from voltage-clamp experiments, are a sufficient explanation of the action potential and of a number of complicated phenomena which have been fitted by solutions of the equations (Hodgkin & Huxley 1952d; Cole 1954; Huxley, unpublished).

\[ V_{Na} \]

\[ V \]

\[ g_{Na} \]

\[ g_{K} \]

\[ V_{K} \]

\[ 150mV \]

\[ 15mV \]

\[ 0 \]

\[ 12mV \]

\[ 4 \]

\[ 2 \]

\[ 0 \]

FIGURE 12. Theoretical action potential and conductance changes obtained by numerical solution of equation (8) and subsidiary equations for \( n, m \) and \( h \); the constants used were appropriate to a temperature of 18.5⁰ C; from Hodgkin & Huxley (1952d). Total entry of \( Na^+ = 4.33 \) pmole/cm². Total exit of \( K^+ = 4.26 \) pmole/cm². Velocity = 18.8 m/sec. Temperature = 18.5⁰ C.

The sequence of events during the nervous impulse

Although the quantitative theory which Huxley and I developed is largely empirical, it gives a fairly clear picture of the sequence of events during the conduction of an impulse in the giant nerve fibre of Loligo.

In figure 12, the horizontal lines give the equilibrium potentials of the sodium and potassium ions. The broken curve is the membrane potential and the two continuous curves are the sodium and potassium conductances. At rest, although the potassium conductance is small, it is much greater than the sodium conductance.
so the membrane potential is fairly close to the equilibrium potential of the potassium ion. As the impulse advances, the membrane just ahead of the active region becomes depolarized by electric currents flowing in a local circuit through the axoplasm and external fluid. Under the influence of the change in membrane potential the sodium conductance rises and sodium ions enter the fibre; this inward current makes the inside of the fibre positive (by carrying charge through the dielectric) and provides the current required to depolarize the resting membrane ahead of the active region. At the crest of the spike the slow changes which result from depolarization begin to take effect. The sodium conductance declines and the potassium conductance rises so that the rate at which potassium ions leave the fibre exceeds the rate at which sodium ions enter; this makes the potential swing towards the equilibrium potential of the potassium ion. As the potential approaches the resting level, any sodium conductance which has not been inactivated is cut off and this may accelerate the rate of repolarization. This effect is conspicuous in myelinated fibres (Tasaki 1956) and although it is not obvious in calculations appropriate to a squid fibre at $18^\circ$ C, those for $6^\circ$ C show it quite clearly (Hodgkin & Huxley 1952a, figure 12). The slow effects of depolarization, namely raised potassium conductance and inactivation of the sodium carrying system, persist for a few milliseconds after the spike and give rise to the refractory period. Under some, but not all, conditions the potential and conductance may return to their resting value in an oscillatory manner. About 10 msec after a spike the membrane is back in its original state and can conduct another impulse. The fibre has gained a small quantity of Na and has lost a similar quantity of K. These movements provide the immediate source of energy for the conduction of impulses and are reversed later by a slow process which requires metabolic energy.

Possible application to other tissues

At present it is uncertain how far the mechanism worked out on the giant axons of cephalopods applies to other excitable tissues, or whether it always applies in these fibres. Hagiwara & Tasaki (1957) have shown that when tetraethylammonium ions are injected into a squid axon, the action potential becomes greatly prolonged and resembles that found in the Purkinje fibres of the heart (Draper & Weidmann 1951). These prolonged action potentials are abolished by removing sodium ions from the external medium, but the ionic properties of membranes altered by injection of tetraethylammonium ions remain to be worked out. From the voltage-clamp records obtained by Hagiwara & Tasaki it would appear that after treatment with tetraethylammonium the rise in potassium conductance, which normally follows depolarization, must either be absent or very greatly delayed.

Since the action potentials and resting potentials of a number of tissues vary with sodium and potassium concentration in approximately the same manner as in the squid fibre, it would seem likely that movements of these ions are involved in many cells. However, there must be some fairly radical differences; thus the action potential of heart muscle lasts about 300 times longer than that in the squid giant axon, and in striated muscle the membrane rectifies direct current in the
opposite direction to that found in nerve (anomalous rectification, Katz 1949). At present it is uncertain whether these differences arise from peculiarities of the systems controlling sodium and potassium permeability, or whether other ions or charged particles are involved. The clearest evidence of propagation in the absence of sodium or of a substitute such as lithium is provided by the experiments of Fatt & Katz (1953) on crustacean muscle fibres. Here it was found that muscles which had previously been treated with tetrabutylammonium ions gave a prolonged heart-like action potential in the absence of all monovalent cations in the external medium. As Fatt & Katz suggested, it would seem that the membrane of a crustacean muscle fibre must either become permeable to one of the internal anions, or perhaps, to calcium and magnesium ions.

Lorente de Nó (1949) showed that tetrabutylammonium ions restore excitability to the B and C fibres of frog nerves after they have previously been made inexcitable by sodium-deficient solutions. Guanidinium and certain other quaternary ammonium compounds will restore excitability to the A fibres of frog nerves which have been blocked with sodium-deficient solutions (Larramendi, Lorente de Nó & Vidal 1956; Lorente de Nó, Vidal & Larramendi 1957). One way of explaining these results is to suppose that certain quaternary ammonium ions substitute directly for sodium ions. However, in view of the results of Fatt & Katz it is conceivable that treatment with quaternary compounds may change the mechanism in such a way that neither sodium nor the quaternary ions is directly involved in carrying current through the membrane. Another puzzling result, described by Müller (1956) is that, after the passage of a large outward current a node of Ranvier will apparently give a prolonged action potential in the complete absence of sodium or potassium from the external medium.

**The nature of the permeability changes**

*Selectivity*

Although little is known about the molecular organization of the membrane there are certain clues as to the nature of the mechanism which controls the permeability to the sodium and potassium ions. In the first place there is evidence that the membrane, under appropriate conditions, is capable of showing a high degree of selectivity towards either sodium or potassium. An example of potassium selectivity is afforded by the variation of membrane potential with potassium concentration during the period of increased conductance which follows the action potential in cephalopod axons. With potassium concentrations above 20 mm, the membrane potential agrees closely with that calculated for a potassium electrode, mean values of $V$ being $-44$ or $-65$ mV with 52 or 21 mm-K in the external solution and a measured internal concentration of 270 mmole-K per litre axoplasm (Hodgkin & Keynes 1955b). With 10 or 3.5 mm-K in the external solution (perhaps slightly more near the membrane) the membrane potential is $-76$ or $-93$ mV, respectively. Since the external solution contained a high concentration of sodium ($[Na]_o + [K]_o = 490$ mm), it is evident that the selectivity of the membrane for potassium is very great. This conclusion can be made more quantitative by
calculating the relative permeability to sodium and potassium by the following equation (cf. Goldman 1943; Sollner, Dray, Grim & Neihof 1955):

$$V = \frac{RT}{F} \ln \left( \frac{[K]_0 + b[Na]_0}{[K]_i + b[Na]_i} \right).$$  \hspace{1cm} (9)

With the data given above, the constant $b$, which expresses the apparent permeability of Na relative to that of K, is found to be about 0.01. Data obtained recently by Horowicz and myself on single muscle fibres immersed in chloride-free sulphate media also give a value for $b$ of about 0.01. The ability of the membrane to discriminate between different alkali metals has not yet been fully worked out, but it seems clear that the membrane is always less permeable to caesium than to potassium. In frog muscle rubidium depolarizes less than potassium (see Keynes & Adrian 1956), but in crab nerve it has a slightly greater effect on both membrane potential and membrane conductance (Wilbrandt 1937; Hodgkin 1947); in Sepia axons, R. H. Adrian found that rubidium depolarized slightly more than potassium. The ability of different tissues to concentrate cations shows a similar variation between K and Rb (Krogh 1946).

During the period of high sodium permeability the discrimination appears to be almost equally great, but the membrane now favours sodium and lithium rather than potassium and rubidium. From the fact that the membrane obeys the equation for a sodium electrode to within 1 mV when the choline concentration is ten times greater than the sodium concentration (Hodgkin & Huxley 1952a, table 1) it follows that the permeability of the active membrane to sodium ions must be at least 200 times greater than that to choline. It is not easy to give a figure for the relative permeability of the active membrane to sodium and potassium, but the voltage-clamp data suggest that the ratio of sodium to potassium conductance reaches at least 30 (Hodgkin & Huxley 1952d). Higher values would probably be obtained if the resting potassium conductance were depressed by hyperpolarizing the fibre before applying the cathodal pulse (Frankenhaeuser & Hodgkin 1957). As was first shown by Overton (1902) the mechanism which produces an action potential does not discriminate between sodium and lithium. Thus, the size of the action potential does not alter to any appreciable extent when sodium ions are replaced with lithium ions (Hodgkin & Katz 1949; Huxley & Stämpfli 1951).

**Variation of conductance with membrane potential**

One interesting point about the system which controls the sodium and potassium conductance of the membrane is that the variation of conductance with membrane potential is extremely steep. Thus an e-fold increase of sodium conductance can be brought about by a change in membrane potential of the order of 5 mV (Hodgkin & Huxley 1952a). In a physical device, such as a vacuum tube or a crystal rectifier, an e-fold change in conductance is usually associated with a potential change of the order of $kT/e$. At room temperature $kT/e$ is 25 mV, and one would expect that a similar quantity would apply to any system in which the conductance was controlled by the movement of particles bearing a single electronic charge. Since the conductance of the membrane changes e-fold in about 5 mV, it would seem that
alterations in permeability must be brought about by the simultaneous movement of a number of charges. These charges might all be located on one particle, or, as mentioned on p. 17, several particles with a smaller charge might be involved at each site where ions go through the membrane.

The quantitative expressions which Huxley and I used to describe the changes in conductance represent a compromise between the ideas of several singly charged particles and one multiply charged particle. Thus the assumption that the potassium conductance is proportional to $n^4$ (p. 17) implies that four particles are involved at each site, while the expressions used for the relation of the rate constants $\alpha_n$ and $\beta_n$ to membrane potential are roughly consistent with each particle being divalent. The evidence (Hodgkin & Keynes 1955b) that potassium ions appear to move in single file, possibly along a chain of sites, may be relevant to the suggestion that several particles are involved at each site where ions go through the membrane.

The influence of calcium

Another line of attack comes from studies of the action of calcium. It has been known for a long time that nerves often become spontaneously active if the concentration of calcium in the external medium is reduced. Recent experiments suggest that the increase of excitability in low calcium depends on an increase of sodium conductance, the curve relating sodium conductance to membrane potential being shifted 6 to 9 mV along the voltage axis for an e-fold increase of calcium concentration (Weidmann 1955; Frankenhaeuser & Hodgkin 1957; Frankenhaeuser 1957). These experiments also suggest that in squid fibres the curves relating potassium conductance and inactivation are shifted in the same direction, though, in the latter case, possibly by a smaller amount. One explanation of the result is that calcium ions are adsorbed on the membrane and that this alters the local electric field inside the membrane without changing the overall potential difference between external and internal solutions. In this case, calcium ions would be important in so far as their concentration influenced the permeability and excitability of the membrane, but they could not be regarded as taking any very direct part in the conduction of impulses. This is perhaps the safest way of interpreting the results, although there are certain facts which point to calcium ions being involved in a more direct way.

One piece of evidence is that a small quantity of calcium enters a squid fibre when it conducts an impulse (Flückiger & Keynes 1955; Hodgkin & Keynes 1957). A resting fibre is only very sparingly permeable to calcium, but the entry can be speeded up about 20-fold by stimulation at 100/sec. The quantity of calcium which normally enters the fibres in one impulse is about 0.006 pmole/cm²; this is only 1/700 of the total sodium which enters the fibre in one impulse and is too small to carry much current through the membrane. The interesting point about the results is that movements of calcium may have something to do with the change in permeability. Since there appear to be substances in the axoplasm with a high affinity for calcium, it is conceivable that depolarization allows calcium ions to be handed on from the membrane to the axoplasm and that this reaction liberates carriers which transport sodium through the membrane.
Movements of ions against concentration differences

Although much more might be said about the action of calcium and the nature of the permeability changes, it is time to consider the fate of the sodium and potassium ions which move across the membrane during the impulse. In giant nerve fibres, the effects of a single impulse are very small; from the figures in table 1 and 2 it can be seen that a 500\(\mu\) axon loses only about one-millionth of its potassium in one impulse. Nevertheless, if a nerve fibre is to be of any use to the animal, it must be equipped with a mechanism for extruding sodium and reabsorbing potassium ions. In contrast to the initial ionic movements during the impulse, the recovery process is slow and, in a large fibre, it may take several hours to wipe out the effects of a short burst of electrical activity.

Sodium extrusion during recovery

The forced ionic movements which occur during recovery may conveniently be studied with radioactive tracers. For sodium, one method (Hodgkin & Keynes 1955a) is to load the fibres with \(^{24}\text{Na}\) by stimulation in a solution containing this isotope. The fibre is then washed with a steady flow of artificial sea water made up with ordinary, inactive sodium. The fluid which flows past the central part of the fibre is collected and its radioactivity measured at intervals. In this way one can study the fate of the sodium ions which enter during activity and are subsequently ejected by the action of a metabolic pump.

Figure 13 illustrates the effect of 0.2 mm-dinitrophenol on the rate at which a Sepia axon eliminates labelled sodium during the period of recovery which follows a burst of electrical activity. At this concentration, dinitrophenol probably does not reduce respiration but it prevents metabolism from doing useful work, possibly by interfering with oxidative phosphorylation. It inhibits movement of ions against concentration gradients in gastric mucosa (Davies 1951), frog skin (Fuhrman 1952), chicken erythrocytes (Maizels 1954) and in certain plant cells (Robertson, Wilkins & Weeks 1951; Scott & Hayward 1954). It does not have any obvious effect on sodium transport in frog muscle (Keynes & Maisel 1954) or mammalian erythrocytes (Maizels 1951), possibly because glycolysis is an important source of energy in these tissues. Applied to a Sepia nerve fibre, it reduces the sodium efflux to about 1/20 of its normal value; the inhibitory effect takes place with a time constant of 15 to 30 min and is largely, but often not completely, reversed by washing the dinitrophenol away. Cyanide (1 to 10 mm) and azide (3 mm) have essentially the same action as dinitrophenol on the outflow of sodium.

Although agents like dinitrophenol or cyanide impair the recovery mechanism, they do not have any rapid effect on the action potential of giant axons, and fibres which have been poisoned with these substances continue to conduct impulses for many hours. The experiment of figure 14 shows that when a squid fibre is treated with dinitrophenol, sodium ions still move into the fibre during activity, but that they accumulate inside the fibre and are not pumped out until the dinitrophenol is removed. The conclusion from experiments of this kind is that the metabolic reactions inhibited by cyanide or dinitrophenol are not of immediate importance.
Figure 13. Action of 2:4 dinitrophenol on the outward movement of sodium in a Sepia axon which had been recovering from the effect of stimulation; from Hodgkin & Keynes (1955a). Temperature 18°C. Abscissa, time after end of stimulation in solution containing $^{24}$Na. Ordinate, rate at which $^{24}$Na leaves fibre.

Figure 14. Effect of 0.2 mm-DNP on sodium entry during stimulation of a squid axon. Temperature 17°C; from Hodgkin & Keynes (1955a). The abscissa is time and the ordinate gives the amount of $^{24}$Na inside the fibre; external $^{24}$Na was washed away with a stream of sea water.
in generating the spike but are essential for recovery. This is very reasonable. During the spike ions move downhill and this does not require metabolic energy—although of course it may need a complicated and tricky mechanism to change the permeability at the right time and in the right way. During recovery, a considerable amount of work has to be done, and it is therefore not surprising that metabolic inhibitors are effective.

Although it may be some time before much is known about chemical events in the membrane, information about the nature of the biochemical reactions which drive the ion transport system can be obtained by the methods used by Caldwell and Keynes. Caldwell (1956) examined the effect of agents which inhibit the sodium pump on the phosphate esters of squid nerve. In a normal fibre, the three main forms of phosphate are inorganic phosphate, arginine phosphate and ATP. When a substance like cyanide or dinitrophenol is applied, at a concentration which reduces sodium efflux, the concentrations of both arginine phosphate and ATP decrease to about one-tenth of their normal value. At the same time there is an increase in inorganic phosphate. The concentrations of ATP and arginine phosphate recover to a considerable extent when the inhibitor is removed and, in general, there is a satisfactory correlation between the concentrations of ATP or arginine phosphate and the rate at which sodium is extruded from the nerve. Caldwell (1957) also showed that if 0.2 mM-dinitrophenol is applied at an alkaline pH, when it has no effect on the sodium pump, it leaves ATP at its normal level but reduces the concentration of arginine phosphate. At a neutral or slightly acid pH, where dinitrophenol reaches a higher concentration in the axoplasm, both ATP and arginine phosphate are reduced, as is the efflux of sodium from the axoplasm. The implication of these results is that dephosphorylation of ATP may be the immediate source of energy for the pump and that arginine phosphate acts by replenishing the ATP. The conclusion is perhaps what might be expected from the analogy of vertebrate muscle, but it is satisfactory to have experimental evidence of a connexion between ATP concentration and sodium transport.

Caldwell & Keynes (1957) have now followed up these results by injecting ATP and arginine phosphate into nerves poisoned with cyanide. They find that injection of ATP partially restores the efflux of sodium, whereas injections of the same solution hydrolyzed by boiling have no effect (figure 15). Arginine phosphate acts in much the same way as ATP, as one would expect from the fact that it probably generates ATP. The effect of both substances is transient, presumably because they are used up in driving the ion transport system, and perhaps in side reactions as well. The quantity of Na extruded was found to be roughly proportional to the amount of ATP injected. On the assumption that all the ATP is used in driving the sodium pump, Caldwell & Keynes (1957) calculate that about four phosphate bonds are broken for each sodium ion expelled. The efficiency calculated from this result, about 10%, seems rather low, but the measurements on which it depends were made in the presence of cyanide and a normal fibre may utilize ATP more effectively. In connexion with these results it should be mentioned that, although an injection such as that used in the experiment of figure 15 raised the ATP concentration in the axoplasm to about the same level as in a fresh fibre, the sodium
efflux was not fully restored. The incompleteness of the restoration might be explained by supposing that ATP is normally produced near the membrane at a higher concentration, or in more general terms, that microinjection of ATP is not a perfect imitation of the normal effect of metabolism. Nevertheless, in spite of uncertainties of the kind usually raised by new results, these experiments support very strongly the idea that high-energy phosphates, such as ATP, are necessary for the process which extrudes sodium from the axon.

![Graph showing efflux of Na over time with ATP injection](image)

**Figure 15.** Effect of injecting ATP on the rate at which labelled sodium leaves a squid axon which has been poisoned with cyanide, from Caldwell & Keynes (1957).

A 780 μ squid axon was loaded by injecting 6700 counts/min of $^{22}$Na over 12 mm. A solution of ATP, which was hydrolyzed by boiling in the first injection but not in the second, was introduced over the same 12 mm length of axon. The solution contained $16 \times 10^{-9}$ mole ATP and should have raised the ATP concentration by about 3 mM.

**Reabsorption of potassium during recovery**

The next question to consider is whether a metabolic system is also required to absorb potassium ions. Here then are two useful kinds of evidence. In the first place, it is found that agents which inhibit or reduce sodium extrusion have a corresponding effect on the uptake of potassium during recovery (Hodgkin & Keynes 1955a). This inhibition takes place without any appreciable change in membrane potential so that it is unlikely to be an electrical consequence of the reduction in sodium transfer. The second point is that when potassium ions are removed from the external fluid the sodium efflux drops suddenly to about one-fourth of its former value and is restored at once when potassium ions are replaced (figure 16 of this article and figure 11 of Hodgkin & Keynes 1955a). Increasing the external potassium concentration five-fold raises the sodium efflux, but the effect is small compared to that of removing potassium.

Perhaps the most obvious explanation of the effect of K-free solution on the outflow of sodium is that the rise in resting potential, associated with removal of potassium, slows up sodium extrusion by increasing the electrical gradient against which these ions have to be forced. This possibility was tested by varying the
membrane potential with an applied current and seeing whether such variations had any effect on sodium efflux (Hodgkin & Keynes 1954). The answer was that large increases in membrane potential—up to 40 mV—had no effect on sodium outflow. Since the rise in membrane potential produced by removing potassium ions from the external fluid was only 5 to 10 mV, it is clear that the effect of K-free solutions on sodium efflux cannot be caused by the change in membrane potential. The most likely possibility is that both the outflow of sodium and the uptake of potassium depend on metabolism, and that the two movements are coupled in a cyclical mechanism of the type shown in figure 17 (p. 32). With such a mechanism one would expect that interfering with metabolism would reduce the uptake of potassium and that removal of external potassium ions might reduce the outflow of sodium.

![Figure 16. Effect of changing the external potassium concentration on the rate at which sodium ions leave a Sepia axon, from an experiment of Hodgkin & Keynes (1955a). The $^{24}$Na was introduced by stimulating the fibre in a radioactive solution at 50/sec for 9 min. The axon diameter was 193 $\mu$m and the temperature 18° C; vertical lines show ±1 s.e.](image)

General nature of the system which moves Na and K against concentration gradients

The tentative conclusion from the experiments mentioned in the previous section is that the same secretory system acts on both sodium and potassium ions and that sodium extrusion may be coupled to potassium influx. Evidence for a coupling between sodium efflux and potassium influx has been obtained in muscle (Steinbach 1952; Keynes 1954), red cells (Harris & Maizels 1951; Shaw 1954; Glynn 1956) and frog skin (Ussing 1954; Koefoed Johnsen & Ussing 1956). An attractive possibility is that there may be a tight coupling between Na and K movements and that one sodium ion is extruded for each potassium ion absorbed. This does not mean that the sodium efflux must be equal to the potassium influx or that the former should fall to zero in the absence of external potassium; for there may be an exchange of sodium ions across the membrane under conditions in which active transport has ceased (see Ussing 1949; Glynn 1956; Swan & Keynes 1956). However, if the idea of a tight coupling is correct it follows that there should be no net movement of sodium against a concentration difference in the absence of external potassium, and that the sodium efflux should fall to zero when both sodium and potassium ions are removed from the external medium. The first prediction is supported by Glynn’s (1956) studies with red cells, but has not been
adequately tested in nerve and muscle. The second appears to be at variance with the observation that an efflux of sodium (which is inhibited by dinitrophenol) can take place from a Sepia axon into an isotonic sugar solution from which all the potassium and sodium have been removed (Hodgkin & Keynes 1955a). However, in such an experiment it is difficult to be sure that the potassium concentration immediately outside the membrane has not been raised by leakage of KCl to an extent which allows a coupled movement of Na and K to continue. Further experiments along these lines are clearly needed; for the time being the position may be summarized by saying that although there is evidence for a loose or partial coupling it has not been proved that the pump operates by ejecting one sodium ion for each potassium ion absorbed.

A physical basis for a coupled system is provided by a hypothesis of the kind discussed by Harris (1954), Shaw (1954), Hodgkin & Keynes (1954) and Glynn (1956). Suppose that potassium enters the cell in association with a carrier (X) and that on the inside of the membrane this carrier is changed chemically into a second molecule (Y) which is selective for sodium and diffuses outward with sodium. On the outside of the membrane the carrier reverts to the potassium form and is once more available for taking potassium into the cell. This system could be interrupted either by removing external potassium ions or by withdrawing the source of energy which is assumed to convert the potassium carrier, X, into the sodium carrier, Y. If the undisassociated compounds KX and NaY were able to cross the membrane much more freely than the dissociated carriers, X and Y, the system would act as an electrically neutral pump which extruded one sodium ion for each potassium ion absorbed.

At present there is no evidence as to the chemical nature of the carriers postulated to explain movements of substances against concentration differences. Even if such carriers can be shown to exist, the problem of chemical identification will not be easy since there are indications that the density of carriers in the membrane must be very low. A few years ago Schatzmann (1953) showed that one of the cardiac glycosides, strophanthasin, inhibited the active transport of sodium and potassium in human red cells. Other cardiac glycosides have since been found to act in a similar manner (Joyce & Weatherall 1955; Kahn & Acheson 1955; Solomon, Gill & Gold 1956; Glynn 1957) and these compounds have been shown to inhibit ion transport in a number of tissues, including muscle (Johnson, 1956; Edwards & Harris 1957) and squid giant axons (Caldwell & Keynes, unpublished). Schatzmann showed that concentrations of strophanthasin which inhibit ion transport do not affect the oxygen consumption or glycolysis of red cells, and it seems that these substances act directly on the membrane rather than on the energy supply to the pump (Glynn 1955, 1957; Whittam 1957). Some of the cardiac glycosides inhibit ion transport at remarkably low concentrations and, from an analysis of the effect of small quantities on red cells, Glynn (1957) concluded that the number of sites at which active transport occurs must be less than 1000 to 2000 per cell. This suggests that the substances concerned with transporting ions through the membrane against concentration differences must be present in very low concentrations indeed. In connexion with Glynn’s results it is interesting
that electron micrographs of the inside of the red cell envelope show small particles about 250 Å in diameter spaced at distances which correspond very roughly to 2000 per cell (Hillier & Hoffman 1953, figure 34, and Hoffman, personal communication).

One interesting point about the active transport mechanism is that it operates with very little time lag. If $^{24}\text{Na}$ is suddenly injected into an axon, these ions start appearing in the external fluid within about 10 sec (Hodgkin & Keynes 1956). Most of this delay is caused by diffusion in the axoplasm and extracellular fluid; there is little time left to allow sodium ions to be collected in, say, a contractile vacuole and to be ejected in the manner in which water is thrown out of an amoeba.

Although lithium and sodium ions appear to be equally effective in generating an action potential, there is evidence that the secretory mechanism may discriminate between the two ions. Thus, lithium appears to be extruded more slowly than sodium from red blood-cells (Flynn & Maizels 1950) and muscle fibres (Swan & Keynes 1956). In the light of subsequent work (Koefoed Johnsen & Ussing 1956), it would seem that the lithium current in frog skin may be due to a passive accumulation of lithium in the skin (Zehm 1954), and that this ion is pumped more slowly than sodium. Ritchie & Straub’s (1957) observations on the after-potentials of mammalian C fibres are also consistent with the idea that lithium ions enter a nerve during activity but are pumped out more slowly than sodium.

*Maintenance of the steady state*

If a nerve or muscle fibre is to remain in a steady state there must be a balance between the rates at which Na and K leak through the membrane under the influence of electrochemical gradients, and the rate at which these ions are pumped by the metabolic transport system. Although little is known about the way in which the internal concentrations of ions are regulated, it is, nevertheless, reasonable to ask whether a neutral pump, which absorbed one potassium ion for each sodium ion ejected, could maintain a cell in a steady state with differences in concentration and electrical potential of the kind observed experimentally. If the pump is strictly neutral it can transfer no charge across the membrane. This means that in the steady state the rate at which sodium ions leak into the fibre must be equal to the rate at which potassium ions leak out. The membrane potential at which this condition is satisfied will depend on the relative permeabilities to Na and K and with certain reservations can be calculated by means of the equation given on p. 22, i.e.

$$ V = \frac{RT}{F} \ln \frac{[K]_o + b[Na]_o}{[K]_i + b[Na]_i} \quad (9) $$

Taking $b$ as 0.01 and using the data in table 1 as a basis for calculation it is found that the resting potential will be about 5 mV less than the equilibrium potential for the potassium ion, i.e. $-70$ mV instead of $-75$ mV. This appears to be consistent with the observations mentioned on p. 6 which indicate that, in a living squid, the resting potential is within a few millivolts of the potassium equilibrium.

* The theoretical arguments in this section were developed in conjunction with Dr P. Horowicz and Dr R. H. Adrian.
potential; the larger difference in isolated axons may occur because branches are cut through in isolating the axon. Experiments with frog's muscle suggest that the resting potential is about 10 mV less than that expected from the potassium concentration ratio, i.e. $-92 \text{ mV}$ instead of $-102 \text{ mV}$ (Adrian 1956).

At present it is considered that while the movements of sodium and to a lesser extent potassium are influenced by the activity of the metabolic transport system, chloride ions distribute themselves passively across the membrane (Conway 1957). In this case the ratio of chloride concentration should be given by the equation

$$ V = \frac{RT}{F} \ln \frac{[\text{Cl}]_i}{[\text{Cl}]_o}. $$

(10)

If this equation is combined with (9) we obtain

$$ \frac{[\text{K}]_o + b[\text{Na}]_o}{[\text{K}]_i + b[\text{Na}]_i} = \frac{[\text{Cl}]_i}{[\text{Cl}]_o}. $$

(11)

Since $b = 0.01$ in muscle this equation approximates to the Donnan equation (12) except when $[\text{K}]_o$ is small compared to $[\text{Na}]_o$:

$$ \frac{[\text{K}]_o}{[\text{K}]_i} = \frac{[\text{Cl}]_i}{[\text{Cl}]_o}. $$

(12)

As is well known from the work of Boyle & Conway (1941), and others, the ratios of the potassium and chloride concentrations in muscle agree satisfactorily with equation (12) provided $[\text{K}]_o$ is greater than about 10 mM. It seems possible that the deviations which occur at lower potassium concentrations are explained satisfactorily by an equation such as (11) which takes account of the imperfect selectivity of the resting membrane towards potassium.

In the theory of Boyle & Conway (1941) the concentrations of $\text{K}^+$ and $\text{Cl}^-$ inside the muscle, and the cell volume were calculated by combining equation (12) with the relations imposed by the conditions of electroneutrality and osmotic balance. A similar argument applies if $\text{Na}^+$ is regarded as a permeable ion which is maintained at a fixed internal concentration $[\text{Na}]_i$ by a neutral pump. In this case, the internal concentrations of K and Cl, and the cell volume, $v$, are determined by equation (11) used in conjunction with

$$ [\text{K}]_i + [\text{Na}]_i + [\text{Cl}]_i + \frac{A}{v} = C \quad \text{(osmotic balance)}, $$

(13)

$$ [\text{K}]_i + [\text{Na}]_i - [\text{Cl}]_i + \frac{zA}{v} = 0 \quad \text{(electro-neutrality)}, $$

(14)

where $C$ is the total concentration of all particles in the external solution, $A$ is the total quantity of indiffusible particles inside the cell and $z$ is the average valency of these particles. It can be shown that if $b = 0.01$ the solutions of equations (11), (13) and (14), which are similar to those obtained from the simpler theory of Boyle & Conway, agree satisfactorily with data obtained on muscle. The conclusion is that, if the permeability to sodium is small compared with that of potassium, a neutral pump, which extruded one Na ion for each K absorbed, could keep a cell in a balanced state with concentrations similar to those in muscle or nerve.
As a by-product, it is interesting to consider what the steady-state concentrations would be in a cell in which the membrane was not selectively permeable to potassium ions. If the permeabilities to Na and K are assumed to be in the same ratio as the mobilities in aqueous solution, the constant $b$ would be 0.67. In this case, if $A$ is taken as a monovalent anion ($z = -1$), solution of equations (11), (13) and (14) gives a result which is similar to that in the human red blood corpuscle. Thus with 5 mm-K, 145 mm-Na and 150 mm-Cl in the external fluid, the internal concentrations calculated on the assumption that $[Na]_i$ is maintained at 15 mm are $[Cl]_i = 106$ mm and $[K]_i = 135$ mm. Such a system resembles the human red blood corpuscle in being able to maintain a steady state with a $[K][Cl]$ product inside the cell about twenty times greater than that in the external fluid.

**Conclusion**

The implication of the experiments considered in this article is that there are two quite distinct systems in a nerve fibre. On the one hand, there is a secretory mechanism which builds up concentration differences by absorbing potassium and ejecting sodium. In parallel with this system are the special channels which allow sodium and potassium ions to move at varying rates down their concentration gradients. There are a number of differences between the two systems. Metabolic inhibitors affect the secretory system but, at all events in giant nerve fibres, seem to have little effect on the system controlling permeability. Changes in membrane potential or changes in external calcium concentration have a large effect on permeability, but do not seem to have much effect on the secretory system. There are also indications of difference in the ability of the two systems to discriminate between the sodium and lithium ions. For these reasons the two systems have been drawn in figure 17 as physically separate. This conclusion may have to be revised when more direct information is forthcoming; on present evidence it seems to be the most satisfactory inference from experiment.

![Diagram](image-url)
Figure 17 provides a convenient opportunity for summarizing the role of sodium and potassium ions in the conduction of nervous impulses. To begin with, the fibre is in its resting condition with a large potential difference across the membrane. The sodium permeability is very low so that only a trickle of ions enters the fibre; this trickle is dealt with by the pump which keeps the fibre in a steady state. When the nerve conducts an impulse the channels open up, allowing first sodium ions and then potassium ions to move down their concentration gradients. These movements generate the action potential and the fibre pays by accumulating sodium and losing potassium. After activity, the original state is restored by the cyclical process which uses metabolic energy to eject sodium and absorb potassium.

The whole mechanism seems admirably designed for a system in which information is conveyed as a series of brief impulses spaced at widely varying intervals. The ionic concentration differences allow electrical energy to be dissipated at a high rate. If the nerve has to handle a large number of impulses in a short time it can run into debt by accumulating sodium and losing potassium. However, sooner or later the debt must be paid off, and this can be done during the periods of rest which alternate with those of more intense activity.

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References


Caldwell, P. C. 1957 The sensitivity to pH of the inhibitory effects of dinitrophenol on squid giant axons. *Biochem. J.* 67, 1P.


Glynn, I. M. 1955 Action of cardiac glycosides on red cells. *J. Physiol.* 128, 56P.


Keynes, R. D. & Lewis, P. R. 1951 The sodium and potassium content of cephalopod nerve fibres. *J. Physiol.* 114, 151.


Whittam, R. 1957 Potassium movements and phosphate metabolism in red cells. *J. Physiol.* 137, 13P.