Absorption of sodium and water by human rectum measured by a dialysis method

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SUMMARY A dialysis method for the study of intestinal absorption is described. Its use has been assessed in animals and normal human subjects and it has been applied to the measurement of rectal transport of sodium and water.

When the luminal solution was of high sodium concentration (145 m-equiv/l), the sodium influx rate (lumen to plasma) was about five times greater than the sodium efflux rate (plasma to lumen). The luminal sodium concentration associated with zero net sodium flux was very low (<15 m-equiv/l). As the mucosa was charged with the luminal side negative, the epithelium must therefore possess a powerful sodium absorbing 'pump'. With isotonic solutions in the lumen, the amount of water absorbed depended on the sodium concentration and when this was 30 m-equiv/l or less, no significant water absorption was detectable. When, however, water absorption was altered by imposing osmotic gradients, sodium absorption was not significantly affected. The luminal solution tended to become isomolar with plasma; osmotic gradients across the epithelium did not develop.

The particular transport properties of rectal epithelium enabling it to remove sodium from the lumen against considerable electrochemical gradients are well adapted to its function.

To study absorption characteristics of intestine in conscious man, perfusion techniques have usually been employed. Among the disadvantages of these techniques is the need to perfuse a relatively long segment in order that the changes of composition produced are large enough to permit accurate measurements. Furthermore, if radionuclides are employed, there may be problems of radiation hazard due to absorption. The dialysis method described in the present paper was devised to overcome some of these difficulties. A tube made of a dialysis membrane and containing solution of a chosen composition is left in contact with the mucosa of an empty segment of intestine for a timed period. The transport properties of the epithelium at that region can then be deduced from changes in composition of the solution. A dialysis method was introduced some time ago by Wrong, Morrison, and Hurst (1961) in order to obtain samples of faecal fluid. Their method involved swallowing dialysis bags, followed by their subsequent recovery from faeces. The method had a different object from that of the present study and so could provide only limited information on the behaviour of the epithelium at any specified region of the intestine. The object of the present work was to study the action of the epithelium at a particular site, and it was therefore most important that the region should be empty and not contaminated with intestinal contents from above or below.

The present report describes the results obtained by applying the technique to examination of the properties of rectal mucosa. Recently, Devroede and Phillips (1970), principally using perfusion methods, concluded that rectal mucosa failed to absorb electrolytes and water. The dialysis method shows clearly that this is not so and that the rectal epithelium absorbs water and possesses a powerful sodium absorbing mechanism capable of removing sodium from the lumen against considerable electrochemical gradients.1

Methods

Investigations were done on two healthy individuals

1A preliminary report on this work was given at meeting of the British Society of Gastroenterology, November 1970.
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and eight patients who had no evidence of bowel disease. The age range of the subjects was 28 to 76 years. Apart from a few minutes necessary to make initial measurements, the subjects continued their ordinary activities during the period of observation. The nature of the procedure was explained to all the subjects. Several normal and sodium-depleted rats, prepared as previously described (Edmonds, 1967), were also studied in preliminary trials.

**Dialysis Tube**
The construction of the tube is shown in Figure 1. A segment of dialysis tube (diameter 0.5 cm, length 8 cm) was knotted at one end and passed over a rubber catheter (diameter 0.2 cm) and secured by a double tie over the rubber sleeve. In a few experiments, the rubber cannula contained an electrode made from polythene tubing filled with agar 4%, KCl 3M.

**Procedure**
The tube was filled using a 5 ml syringe before the dialysis tube was tied. Air bubbles were expelled and the tube was tied in position, a glass spigot replacing the syringe. The tube was checked for leaks and weighed. It was inserted into the rectum at sigmoidoscopy on the first occasion but subsequently it was placed blindly after lubricating the tip with KY2 water-soluble jelly. The tube was inserted so that it was lying in the rectum with its base about 2 cm from the anus. The tube caused no discomfort and did not interfere with normal activities. It was removed by gentle traction, usually after one hour. Specimens or tubes showing faecal contamination were discarded. In some experiments, the rectum was subsequently rinsed twice with 20 ml saline using a rubber catheter and syringe, the fluid being withdrawn and collected for measurement of radioactivity. In the animal studies, experiments were done on the descending colon, exposure and cannulation being as previously described (Edmonds, 1967). A 4 cm-length of dialysis tube was used and absorption measured both when solutions were in the tube (the distal cannula being removed for this purpose) and when solutions were placed for timed periods in the cannulated segment.

**Measurements of Potential Difference**
Studies in man were done using probe and reference electrodes and a millivoltmeter of the type described by Edmonds and Cronquist (1970) except that KCl 3M was used instead of NaCl 150 mM in the electrodes. The reference electrode was placed on abraded skin of the ventral surface of the forearm. The measurements were made at the end of the exposure period, 10 ml of solution similar to that used in the tube being first injected into the rectum. In a few experiments, measurements were made using an electrode contained within the dialysis tube as shown in Figure 1. In the rats, measurements of potential difference were made as previously described (Edmonds, 1967).

**Chemical and Radionuclide Methods**
Solutions placed in the tube were prepared from stock solutions of NaCl 150 mM, KCl 150 mM, and mannitol 500 mM. Sodium and potassium were added to some solutions, the radionuclide content not exceeding a total of 5 nCi/ml except in a few animal experiments where up to 200 nCi/ml was used. Specimens collected were frozen for storage where analysis was delayed. Water loss was determined by weighing, sodium content measured by flame photometry, and osmolality by a Fiske osmometer. Changes of volume of the fluid within the tube were determined by weighing before and after each experiment. Sodium and potassium were determined in 1 ml samples using a well type NaI (Th) crystal scintillation counter; where both radionuclides were used, the samples were recounted after allowing several days for potassium to decay. The content of potassium in the colon mucosa of the rat was measured in vivo using a miniature Geiger counter (Barnaby and Edmonds, 1969).

**Theory and Calculations**
It is supposed that when the tube is in an empty segment of intestine it is surrounded by a thin layer of fluid which also lies in contact with the epithelium. Ions, water, and other substances can then diffuse from the fluid in the tube through the membrane and across the thin fluid layer, and are then transported across the epithelium into the blood. Flows of ions, water, and other substances will also occur in the opposite direction. In this system, the thin intermediate fluid layer is assumed to behave as a compartment of small capacity. Providing that the
dialysis membrane is very permeable to the substance under investigation and the epithelium is consider-
ably less permeable to it, the concentration of the substance in the intermediate fluid layer will rapidly
approximate to its concentration in the tube. Once
this steady state has been attained, the presence of
the dialysis membrane can be ignored and the
change in the amount of substance in the tube will
depend on the rate at which it passes from the
intermediate layer into the plasma. To measure the
flux rates in the present experiments, sodium
was contained in the solutions in the tube and
calculations were based on the assumption of one
compartment (the luminal solution) and a large sink
(the body), which seems to be adequate even for
potassium calculations (Barnaby and Edmonds,
1969). Since changes of specific activity were
relatively small, it was assumed that the fall with
time was linear during exposure and accordingly
the arithmetic mean was used in the calculations.
Equations used were:

\[
J_l = \frac{V_o C_o - V_i C_i}{\text{s.a.} \times A \times t}, \quad \text{(1)}
\]

\[
J_n = \frac{V_{o,0} - V_{i,t}}{A \times t}, \quad \text{(2)}
\]

\[
J_o = J_n - J_i, \quad \text{(3)}
\]

where \(J_l\), \(J_n\), and \(J_o\) are influx (lumen to plasma),
et flux, and efflux (plasma to lumen) rates per
min : cm² respectively; \(V\), \(C\), and \(I\) are volume of
tube fluid, radionuclide content (cpm/ml) and
centration of the ion; \(\text{s.a.}\), the mean specific
activity; \(A\), the surface area of the dialysis tube;
and \(t\), the time of exposure. The subscripts, \(o\) and \(t\),
refer to initial and final values. The conventional
sign usage is employed, influx (insorption) being
positive and efflux (exsorption) being negative. All
flux and absorption rates are in terms of the area
(cm²) of the surface of the dialysis tube.

Results

ASSessment of the Method

A number of preliminary studies in normal subjects
and in rats were carried out and the results will be
briefly summarized. When a solution of the same
composition was used several times during one day
in two subjects, it was shown that the results were
reproducible (Table I). The greatest variation was of
weight change (SD ± 25%) while the sodium and
potassium concentrations had a standard deviation
not greater than ± 15%. For good consistency, it
was, however, most important that the rectum was empty. The presence of faeces was obvious from
brown staining of the tube fluid and such tubes
were always discarded.

In experiments on rats, changes in composition of
the luminal solution, whether placed freely in the
lumen or within a dialysis bag, were examined. The
results showed that the presence of the dialysis
membrane had no significant effect on the magnitude
of the changes observed (Table II). Also, both in
rats and in human subjects, measurements of potential difference made by the electrode within
the dialysis tube were compared with those made
without the tube. In all instances, no significant
differences were observed. The dialysis membrane
does not, therefore, introduce artifacts into the
measurements of potential difference.

<table>
<thead>
<tr>
<th>Concentrations (m-equiv/l)</th>
<th>Sodium (Cpm/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial solution</td>
<td>30</td>
</tr>
<tr>
<td>After one hour</td>
<td>29 ± 0.8</td>
</tr>
<tr>
<td>No tube</td>
<td>31 ± 0.7</td>
</tr>
<tr>
<td>Solution within tube</td>
<td>505 ± 9</td>
</tr>
</tbody>
</table>

Table II Comparison of changes in composition of a luminal solution in the rat descending colon

1In some experiments the solution was free in the lumen and in others contained in dialysis tube (mean ± 1 SD). Results were obtained from two rats with a total of four observations in each case.

In some experiments, the changes in composition of the tube fluid were compared when the same
solution was exposed with the tube in the rectum,
in a stirred bath of saline, or in a flat, impermeable
polythene bag. In the latter condition, the surface
of the tube was in contact with the polythene. The
sodium content fell much less when the tube
was in the rectum than when it was in a saline bath
(Fig. 2) indicating that the sodium permeability of
rectal mucosa was much less than that of the dial-
ysis membrane, a necessary condition for the flux
calculations. The weight loss was, however, much
greater when the tube was in the rectum. No
significant change was found when the tube was in
the saline and only a very slight fall when it was in
the polythene bag. In the latter case, at the end of

<table>
<thead>
<tr>
<th>Weight Loss of Tube (mg)</th>
<th>Concentrations (m-equiv/l)</th>
<th>Potassium (Cpm/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial solution</td>
<td>90</td>
<td>50</td>
</tr>
</tbody>
</table>
| After one hour
| Subject 1               | 0.21 ± 0.05                | 61 ± 8.5          | 53 ± 3.6          | 340 ± 20          |
| Subject 2               | 0.19 ± 0.05                | 74 ± 3.4          | 48 ± 4.4          | 460 ± 28          |

Table I Consistency of changes in luminal solution during repeated periods of exposure of one hour of the dialysis tube in the rectum of two subjects (mean ± 1 SD)₁

₁Four observations were made in each subject.
the hour, a thin film of fluid was evident over the area of contact of polythene and tube.

On four occasions after removal of the tube, following an hour's exposure, the rectum was rinsed twice with 10 ml saline and the washings were collected and the radionuclide content was measured. The amount recovered from the lumen in this way was 5-9% of the total amount lost from the tube. Thus, during the exposure of an hour, at least 90% of that lost from the tube is absorbed.

Experiments on three rats were done to estimate the extent to which ions spread beyond the limits of the tube along the surface of mucosa. $^{42}$Potassium was added to the tube fluid since this is temporarily 'trapped' in the epithelial cells and can easily be measured (Barnaby and Edmonds, 1969), and so can act as a marker to indicate the spread. The $^{42}$Potassium content of the epithelium was maximal over the region where the epithelium was in direct contact with the tube (Fig. 3) and, although within 1 centimetre of the ends of the tube, the count rate was still about 50% of the maximal value; outside this limit the epithelium contained very little radioactivity. It seems likely, therefore, that nearly all the ionic movements between the luminal solution and the plasma take place through the epithelium in close contact with the tube surface.

**RECTAL TRANSPORT OF SODIUM AND WATER**

**Sodium**

In studies in two subjects, a series of tests of exposure over one hour were done, each successive solution being prepared so as to have the same sodium concentration as that found at the end of the previous exposure. Both subjects showed similar changes and the changes in one of these are shown in Figure 4. Sodium concentration fell until it was

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**Fig. 2.** Changes of weight and $^{22}$Sodium concentration of tube fluid (NaCl 145 mM) left for one hour under various conditions. The $^{22}$Sodium changes are plotted on a logarithmic scale. ● = tube in rectum; ○ = tube in stirred saline bath; □ = tube in a polythene bag.

**Fig. 3.** $^{42}$Potassium content of mucosal epithelium of rat descending colon after exposure for one hour, the dialysis tube containing a saline solution and $^{42}$Potassium. Regions of colon beyond 1 cm from the end of the tube contained little $^{42}$Potassium. The position of the tube is shown.

**Fig. 4.** Changes in sodium concentration of solutions in dialysis tubes during successive one-hour exposures. At the end of each exposure the final sodium concentration was determined and a solution of this concentration prepared (with mannitol added to ensure isotonicity) for the subsequent exposure.
about 10 m-equiv/l. When, however, a solution of sodium concentration 3 m-equiv/l was placed in the rectum, its sodium concentration rose. With solutions of higher sodium concentration, water absorption from the tube, as indicated by a fall in weight, always occurred but when the sodium concentration was very low, no absorption was detectable. In several subjects, a solution of NaCl 145 mM, KCl 5 mM with \(^{22}\)Sodium added was used and the influx (lumen to plasma) determined (Table III). Although the content of \(^{22}\)Sodium fell in these experiments (Fig. 2), so also did the concentration of sodium so that changes of specific activity were insignificant. This indicated either that the net flux was zero or that it was relatively small compared with the influx so that when the amount of sodium present in the tube was high, the effect of the efflux was masked. Hence, in order to determine the sodium flux rates in most subjects, a solution of lower sodium concentration (30 m-equiv/l) was used (Table III). Even so, influx exceeded efflux so that in all subjects, the luminal sodium concentration fell during the exposure, the final value averaging 15 m-equiv/l (range 10 to 19 m-equiv/l). Thus, for the net sodium flux rate to be zero, the luminal sodium concentration must generally be about 15 m-equiv/l.

![Graph](http://gut.bmj.com/)

**Fig. 5. Flux rates of sodium in a normal subject with solutions of various sodium concentrations in the lumen.** Solutions were prepared with varying NaCl concentrations and constant KCl concentration at 30 mM with mannitol added to make solutions isotonic. The values are plotted against the mean sodium concentration during the exposure. ● = sodium influx rate (J1); ○ = sodium efflux rate (J0).

Table III Sodium unidirectional flux rates measured in subjects with normal rectum (mean ± SD).^1^  

<table>
<thead>
<tr>
<th>Initial Sodium Concentration (m-equiv/l)</th>
<th>No. of Subjects</th>
<th>Influx (n-equiv/min/cm²)</th>
<th>Efflux (n-equiv/min/cm²)</th>
<th>Net Flux (n-equiv/min/cm²)</th>
<th>Potential Difference (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>145</td>
<td>4</td>
<td>112 ± 23</td>
<td>—</td>
<td>112 ± 23</td>
<td>-25</td>
</tr>
<tr>
<td>30</td>
<td>8</td>
<td>48 ± 14</td>
<td>-24 ± 9</td>
<td>23 ± 10</td>
<td>(-19 to -32)</td>
</tr>
</tbody>
</table>

^1^The negative sign on a flux rate indicates that the flow was directed towards the lumen. The higher and lower sodium concentration solutions were prepared with potassium concentrations of 5 m-equiv/l and 30 m-equiv/l respectively. It was not possible to measure efflux rate when the luminal sodium concentration was 145 m-equiv/l.

In one subject, the unidirectional flux rates were determined using a range of solutions of various sodium concentrations with \(^{22}\)Sodium; potassium concentration was constant at 30 m-equiv/l (Fig. 5). Estimating the net flux when luminal sodium concentration exceeded 50 m-equiv/l was unsatisfactory but the results obtained at lower concentrations suggested that the net flux did not vary significantly with increasing luminal sodium concentration. The influx increased linearly with increasing luminal sodium concentration over the range studied without any evidence of saturation kinetics. The potential difference measured at the time of the flux determinations (the luminal sodium concentration being about 30 m-equiv/l) averaged −26 mV (range −16 to −33 mV). There was some dependence of potential difference upon sodium concentration but the effect was small; for example, in one subject the potential difference averaged −24 mV with 150 mM NaCl in the tube and −19 mV when it contained 10 mM NaCl. These small changes of potential difference are of similar magnitude to those observed by Dalmark (1970) when the sodium concentration of solutions placed in the rectum was reduced.

**Water**

When the luminal solution had an initial sodium concentration of 145 m-equiv/l, the average weight loss of the tube was 0.22 ± 0.04 g (four subjects, mean ± SD) corresponding to an average water absorption rate of 0.24 μl/min/cm². When the luminal solution had a low sodium concentration (initially 30 m-equiv/l) the average weight loss was much less at 0.05 ± 0.04 g (eight subjects, mean ± SD), a value not significantly different from zero. In one subject, a number of exposures using solutions of high or low sodium concentration were carried
out, all solutions being made isotonic by addition of mannitol, and the results clearly confirmed those above in showing that water absorption rate was greater when luminal sodium concentration and sodium absorption rate were higher (Fig. 6a). To see whether changes in the rate of water flow affected sodium absorption, hypotonic and hypertonic solutions having similar sodium concentrations were used. Thus water activity was varied but electrolyte activity was not. This resulted in considerable change in net water movement from absorption to secretion but sodium absorption was unaffected (Fig. 6b). These results are consistent with the view that water absorption from isotonic solutions by the rectum depends on active absorption of sodium probably by development of local osmotic gradients within the tissue as suggested for other epithelia (Curran and McIntosh, 1962; Diamond and Bossett, 1967).

In several of the experiments in which isotonic solutions of various sodium concentrations were used, the osmolalities were determined after exposure. The average osmolality was 298 ± 12 mOsm (16 observations, mean ± SD) which did not significantly differ from the initial value. Also, when in some trials hypotonic solutions (about 200 mOsm) were used, the osmolality was invariably greater at the end of one hour exposure (average value 242 mOsm, five experiments) while if the solution was initially hypertonic (about 400 mOsm), the osmolality fell (average final value 354 mOsm, three experiments). The potential difference was not significantly affected by changes in osmolality of the luminal solution.

There was no evidence in these experiments that significant osmotic gradients could develop across the rectal epithelium and therefore when sodium concentration fell, its place must have been taken by other substances. Potassium appears to be one of those which replaces sodium but others may also be secreted into the lumen.

**Discussion**

Several of the assumptions in the theory of the dialysis method which were tested in the preliminary assessment appeared to be true. For example, the model experiment with the tube in a polythene bag demonstrated the thin fluid layer that was present between the tube and an adjacent membrane. The weight loss of the tube in this experiment indicated the small volume of the layer, an observation consistent with the finding that little radioactivity was recoverable from the rectal lumen after an exposure. The animal experiments indicated that the presence of the dialysis membrane made little difference to the estimate of ionic movements and potential difference, and also that nearly all the movements from the tube to the animal took place over the area of contact. Thus, in expressing the results in relation to the area of the tube surface, they should approximate to the area (cm²) of mucosal surface. The principal advantage of the present technique is that a short defined segment of intestine can be studied by a sensitive method. Although the rectum is the most obvious and easy segment to investigate, because it is accessible and usually empty, with some modifications possibly other parts
of intestine could also be examined. The method is sensitive because small volumes of solution can be held at a chosen site for fairly long periods, a procedure which is otherwise not practicable in conscious man. Furthermore, it is an advantage that if radioactive tracers are used, very little activity is absorbed.

The particular sensitivity of the dialysis method is demonstrated by the results of the rectal studies. Employing perfusion methods, Devroede and Phillips (1970) were unable to show significant sodium or water absorption and concluded that, in contrast with the rest of the colon, the rectum ‘failed to absorb water and electrolytes’. This result was surprising as rectal mucosa is electrically polarized like the rest of the colon (Geall, Spencer, and Phillips, 1969; Dalmark, 1970) and the potential difference rises considerably when aldosterone or other mineralocorticoids are administered (Edmonds and Godfrey, 1970). Similar changes in rat colon are associated with increased sodium absorption (Edmonds and Marriott, 1967) so that the rise of potential difference in man suggested the presence of a sodium absorptive mechanism in rectal mucosa. The dialysis method gave results consistent with this suggestion demonstrating clearly that rectal mucosa is not inert in regard to the transport of ions but can remove sodium from the lumen against considerable electrical and sodium concentration gradients. The epithelium must, therefore, possess a highly efficient active sodium transport mechanism.

It is difficult to make satisfactory comparisons with absolute flux rates obtained by perfusing the whole colon. Nevertheless, the present results are of similar order to those obtained by perfusion studies, although showing some differences. Thus Shields (1966) measured flux rates using a luminal solution (Tyrode’s) with sodium and potassium concentrations of about 150 and 5 m-equiv/l respectively. If for approximate comparison they are expressed in cm², the colon being treated as a tube 150 cm long and of 4 cm in diameter, the average values obtained for sodium were: influx = 338, efflux = 151, expressed in n-equiv/min/cm². The present results obtained with a solution of similar sodium concentration in the rectum gave a value for influx of only 112 n-equiv/min/cm², considerably less than for the colon as a whole. Furthermore, if the relative values of efflux and influx are considered, a striking difference is evident. For whole colon, the efflux had a value of about 45% of the influx, but for the rectum efflux was only about 20% of influx (assuming the value of efflux shown in Table III). Sodium efflux is probably passive and these findings, therefore, strongly suggest that permeability to passive sodium movement is considerably less for rectal epithelium than for that of other parts of the colon.

The characteristics of colonic epithelium probably differ in various regions. For example, the response to aldosterone is much greater in the distal than in the proximal colon (Edmonds and Marriott, 1967). It would be expected that the variations reflect functional requirements. An important function of distal colon and rectum is sodium conservation and sodium has to be removed against considerable electrochemical gradients as here the faecal fluid has low sodium concentration. The present results, indicating the restricted permeability of rectal mucosa to passive ionic diffusion together with the presence in the epithelium of a powerful sodium absorbing ‘pump’ would seem well suited to the functional requirements of this region of intestine.

References


