Water and sodium absorption by the intestine in cholera

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How the copious diarrhoea in acute cholera is produced is poorly understood. It may be due to overproduction of fluid normally entering the gastro-intestinal tract, to failure of reabsorption of the normal volume of such fluid, or to a combination of both defects (Watten, Morgan, Songkha, Vanikiati, and Phillips, 1959). The demonstration of an inhibitor of sodium transport, using short-circuited frog skin, in cholera stool has focused attention recently on the major role of a defect in absorption rather than on overproduction (Huber and Phillips, 1960).

The present studies investigated the changes in water and sodium transport in the rabbit small intestine which had been shown (De and Chatterje, 1953) to accumulate luminal fluid when exposed to vibrio or their products. This animal model may more closely resemble the human intestinal disturbances in cholera than frog skin preparations.

METHODS

Rabbits of local breed, weighing 1·5 to 2·kg, were anaesthetized by intravenous nembutal. Isolated loops of jejunum, 8 to 10 cm in length, were constructed, leaving their blood supply intact on the mesenteric pedicle. The ends of the loops were cannulated to allow fluids to be introduced and removed. The continuity of the remaining small intestine was reestablished. The loops were replaced in the abdominal cavity and the midline incision was closed around the cannulae. Two loops were prepared in each animal. Studies either began immediately in the case of experiments with the animals exposed to toxins or an inoculum of vibrio was made into one of the loops to be studied at intervals subsequently. A freshly prepared culture of vibrio was placed in one of the two loops. A similar volume of the same material heated to 80°C for 30 minutes was placed in the other loop. The cephalic and caudal loops were used alternately as control and experimental loops. In all cases the cholera vibrio was an Inaba strain (NIH type 35A3) harvested after two hours’ incubation in alkaline peptone water at 37°C. The inoculum consisted of 10^7 to 10^8 organisms suspended in 1 ml saline. In other studies the experimental loop was exposed to cholera toxin (Jenkin and Rowley, 1959) produced by ammonium sulphate precipitation of sonicated vibrios. Cholera filtrate (Huber and Phillips, 1960) was also studied after Seitz-filtering stools from two cholera patients; in both these cases 1 ml of toxin or filtrate containing 1 mg protein per millilitre was inoculated. Twenty-six studies were carried out with vibrio infection and 12 each for both toxin and filtrate.

To study water and sodium transport in the loops 5 ml of an electrolyte solution containing sodium 140, potassium 10, chloride 110, and bicarbonate 40 m-equiv/l. was placed in the intestinal lumen for 10-minute periods. The solutions were injected at body temperature which was maintained at 37°C throughout the studies. Net absorption was calculated from input-output balance, estimated by volumetric recovery and in some studies by changes in dilution of a nonabsorbable marker (PEG). Chemical determinations of sodium and potassium were made with a Coleman flame photometer, of chloride by a Cotlove titrator, and of bicarbonate as CO₂ by the manometric method of van Slyke. Osmolalities were measured with a Fiske osmometer.

Unidirectional fluxes of sodium ion were determined simultaneously by using both isotopes of sodium. The extracellular fluid of each animal was labelled by injecting 2 microcuries of ^24Na intravenously. This was done at least two hours before absorption studies commenced to allow adequate mixing. The electrolyte solution instilled into the loops was labelled with ^22Na at a concentration of 2·5 microcuries per litre. Sodium 22 and 24 activities were determined by counting loop fluid samples immediately after the study and then recounting after a lapse of one week when the levels of sodium 24 had decayed to negligible values. Counting was carried out in a 2 × 2 sodium iodide well-type scintillation counter, 2 ml samples being counted to a minimum of 10,000 counts. The unidirectional fluxes were calculated as follows:

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\text{Lumen to plasma (L-P)} = \left( \frac{\text{total } Na \text{ m-equiv}}{\text{injected}} \right) \left( \frac{\text{Counts/min injected} - \text{counts/min recovered} \times 100}{\text{Counts/min injected}} \right)
\]

\[
\text{Plasma to lumen (P-L)} = \frac{\text{Specific activity of } ^24Na \text{ in plasma (CPM/m-equiv)}}{\text{Counts/min recovered}}
\]
These values are expressed as \( \mu \)-equiv (sodium ion)/g wet intestine weight/hour.

**RESULTS**

Figure 1 shows the net transport of water and electrolytes across the intestinal mucosa in control and vibrio-infected loops. In the normal loop net absorption was 2.32 ± 0.40 ml/g/hr whereas in the infected loop net accumulation 1.48 ± 0.32 ml/g/hr was observed. The values are the mean and standard deviation of 26 paired studies. Similar changes were seen in net sodium transport and if the composition of fluid either absorbed or accumulating is calculated it is approximately the same concentration of sodium ion as plasma, 140 m-equiv/l. Chloride and bicarbonate, which were absorbed by the normal mucosa, accumulated in the infected loop. Little change was seen with potassium transport.

![Fig. 1. Net transport of water and electrolytes in normal and infected intestinal loops. Vibro infection 12 hours.](image)

The plasma flux and plasma-to-lumen flux directly. In these latter studies there was close agreement between the net transport measured chemically and algebraic summation of the separate ionic fluxes. In Fig. 3 the effect of time of exposure to the vibrio on sodium ion fluxes is shown. After four hours' infection absorption still occurred but was less than in the control; at eight hours net accumulation in the lumen of sodium began and this was accentuated after 12 hours. After 24 hours of infection the bowel behaved as an inert membrane with no net movement across the mucosa. Exposure to cholera toxin produced a more rapid effect.

![Fig. 2. Net water transport, net sodium, and unidirectional ion fluxes in control, vibrio-infected, J.R. toxin, and cholera filtrate exposed intestinal loops. Infection time 12 hours. Toxin exposure time four hours.](image)

![Fig. 3. The effect of length of exposure of the intestine to cholera vibrio on net water, net sodium absorption, and unidirectional sodium ion fluxes in intestinal loops.](image)
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which reached a plateau of activity after one hour (Fig. 4).

The transport of water and sodium by mucosal cells of both the jejunum and ileum was disturbed by exposure to vibrio. The main difference is that the unidirectional fluxes of sodium ion were greater in the jejunum (Fig. 5). The changes in net transport of sodium in the infected loop were due to an increase in plasma-to-lumen flux in both areas of the small intestine. These findings are in agreement with those of Fordtran, Rector, Ewton, Soter, and Kinney (1965) who have demonstrated greater permeability of the jejunum than of the ileum in man.

Figure 6 shows the effects of glucose absorption on water and sodium transport in the normal infected loops. Glucose was present in the luminal fluid at a concentration of 20 mMol. Simultaneous absorption of glucose enhanced the lumen-to-plasma flux of sodium ions by 29.4%. This resulted in a 46.4% increase in net sodium absorption. Net water absorption is similarly increased by 17.5%, the fluid transported remaining isotonic. There was no change in the plasma-to-lumen sodium flux. In the infected loop glucose absorption was not decreased, being $84.4 \pm 10.8 \mu\text{mol/g/hr}$ in the controls and $82.3 \pm 12.2 \mu\text{mol/g/hr}$ in the infected loops. Glucose absorption increased the sodium flux from lumen to plasma to a similar degree to that in the control series. This enhancement of sodium ion flux resulted in a change from net accumulation in the lumen of $204.3 \pm 48.5 \mu\text{equiv/g/hr}$ to net absorption of $102.8 \pm 26.4 \mu\text{equiv/g/hr}$ in the infected loop. Water was absorbed in isotonic proportion, net absorption being produced by glucose transport. There was no change in the plasma-to-lumen flux of sodium in the infected loop during simultaneous absorption of glucose.

**DISCUSSION**

The main symptom in cholera infection is the rapid loss of a large amount of fluid by the intestinal wall. Contrary to an earlier view, this occurs while the patient retains a morphologically intact intestinal mucous membrane. It therefore appears that the vibrio is causing a subtler change in the mucosal epithelium than its removal. A possible basis for
understanding this change lies in the unidirectional fluxes of water and sodium ions across the intestinal mucosa (Visscher, Vargo, Carr, Dean, and Erickson, 1944; Fordtran, Levitan, Bikerman, Burrows, and Ingelfinger, 1961), their magnitude in man (Love, Mitchell, and Phillips, 1968) being sufficient to account for the large volumes of diarrhoeal stool found in the human disease.

The present studies using a loop of adult rabbit small intestine in vivo have shown that the loop becomes distended with fluid having the composition of cholera stool. This was due to an increase in the unidirectional flux of sodium ion from plasma to lumen and so to a net accumulation of sodium in the intestinal lumen. Huber and Phillips (1960), however, believe that the effect of the vibrio is to decrease the flux of sodium ion from lumen to plasma. This conclusion is based on the observation that vibrio inhibit the short circuit current of the frog skin. The short circuit current in the frog skin and small intestine (Schultz and Zalnsky, 1964) is a measure of net active sodium transport and not of unidirectional flux. It will be decreased, therefore, by either reduction in lumen-to-plasma flux or an increase in plasma-to-lumen flux (Fuhrman, 1952). The results of Huber and Phillips (1960) could therefore be equally well explained by an increased movement of sodium ions into the intestinal lumen rather than by decreased lumen-to-plasma movement. This would be in agreement with the evidence now presented from the small intestine.

This concept of increased movement of sodium ions through the mucosal cell is also in keeping with other work. Ling (1965) believes that the term 'sodium pump' does not imply anything more than a cellular barrier to massive fluid and electrolyte movement by normal mucosa. In cholera this barrier appears to be disrupted so that the cell cannot accumulate sodium against a concentration gradient. This leakiness of mucosal cells may not even be transcellular but involve changes in the 'shunt path' (Ussing and Windhager, 1964). Cholera stool and vibrio filtrates have been shown to affect the permeability of capillaries in the skin of guinea pigs and rabbits (Craig, 1965) so that the observed effect in the intestine may not be peculiar to intestinal epithelium.

The accumulation of fluid in intestinal loops has been demonstrated in infected bowel where a number of toxic vibrio products may be operating. In the studies exposing the mucosa to toxin without vibrio the results were similar but must be interpreted with caution, since in the preparation of the Jenkin-Rowley toxin a concentration of 0·8 mM ammonium chloride is finally present in the final product. This concentration of ammonium chloride has been shown to inhibit fluid absorption in intestinal sacks and loops and also to reduce the short circuit current produced by active sodium transport by the frog skin (Neptune and Mitchell, 1965).

Further evidence of the integrity of the lumen-to-plasma-sodium flux is seen in the absorption of glucose by the infected loops. Czaky (1964) believes that glucose and sodium share a common transport system. In the present studies glucose absorption was not depressed in the cholera loop, and if the common transport hypothesis is correct, then this would make the isolated inhibition of the sodium pump unlikely. These absorptive processes are thought to be dependent on ATPase (Newey, 1967). Studies by Richardson and Evans (1965) have shown that vibrio extracts have no effect on the ATPase activity in preparations of intestinal microvilli. It is possible that some of the glucose effect is due to solvent drag, but, according to Ussing (1963), this would decrease the plasma-to-lumen flux which has not been observed in the present studies. The finding that simultaneous glucose absorption enhances lumen-to-plasma flux of sodium, and consequently net sodium and water absorption in the presence of cholera vibrio is further evidence of the metabolic integrity of the mucosal cell.

The present studies therefore indicate that the accumulation of fluid in the cholera-infected intestine of the rabbit is due to excessive movement of sodium ions into the lumen and not to depression of ionic movement out of the lumen. This seems to represent increased cellular permeability rather than inhibition of the mucosal sodium pump. This conclusion is in agreement with the original suggestion of Becquerel (1849).

**SUMMARY**

The effects of cholera vibrio and their products have been studied in vivo in a rabbit intestinal loop preparation. The accumulation of luminal fluid similar to cholera stool in man was observed. The net sodium content was accounted for by an increase in plasma-to-lumen unidirectional ionic flux and not to a decreased lumen-to-plasma flux. Simultaneous glucose absorption enhanced the latter flux and could result in net sodium absorption. These findings are consistent with an increased mucosal cell permeability rather than sodium pump inhibition.

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