Ca²⁺ Transport across Intestinal Brush Border Membranes of the Cichlid Teleost Oreochromis mossambicus

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Summary. Brush border membranes were isolated from tilapia (Oreochromis mossambicus) intestine by the use of magnesium precipitation and differential centrifugation. The membrane preparation was enriched 17-fold in alkaline phosphatase. The membranes were 99% right-side-out oriented as indicated by the unmasking of latent glyceraldehyde-3-phosphate dehydrogenase and acetylcholine esterase activity by detergent treatment.

The transport of Ca²⁺ in brush border membrane vesicles was analyzed. A saturable and a nonsaturable component in the uptake of Ca²⁺ was resolved. The saturable component is characterized by a \( K_m \) much lower than the Ca²⁺ concentrations predicted to occur in the intestinal lumen. The nonsaturable component displays a Ca²⁺ permeability too high to be explained by simple diffusion. We discuss the role of the saturable component as the rate-limiting step in transmembrane Ca²⁺ movement, and suggest that the nonsaturable component reflects a transport mechanism operating well below its level of saturation.

Key Words  Ca²⁺ transport • brush border membrane • enterocyte • fish intestine • tilapia

Introduction

In freshwater fish, the gills are the major site for Ca²⁺ uptake from the (hypocalcic) water environment [8, 41]. In combined in vivo and in vitro studies Flik et al. [10, 12, 13] identified a high-affinity Ca²⁺-ATPase as the pivotal mechanism in active Ca²⁺ transport in the gills of freshwater tilapia (Oreochromis mossambicus Peters) and North-American eel (Anguilla rostrata LeSueur).

Uptake of calcium via the intestine provides a second important pathway. Indeed, Berg [1], Dacke [4] and Ichii and Mugiya [22] described important contributions of dietary calcium to the total Ca²⁺ uptake in fish. The prevailing intracellular Ca²⁺ concentrations (≈80 nmol · l⁻¹) are nonsaturating for the basolateral Ca²⁺ extruding mechanisms [11, 34]. The rate-limiting step of transepithelial Ca²⁺ transport therefore seems to be the Ca²⁺ entry at the apical membrane. The primary site of hormonal control of Ca²⁺ uptake has been suggested to be in the apical membrane of cells in the gills and intestine [25, 30]. In tilapia, transepithelial Ca²⁺ transport is sodium dependent and associated with a powerful Na⁺/Ca²⁺ exchanger. Furthermore, the enterocyte is equipped with a high-affinity ATP-dependent Ca²⁺ transporter [11, 34].

In fish, at least in some species, the calcium metabolism is hormonally controlled by prolactin or cortisol that exert hypercalcemic effects, and by the hypocalcemic hormone stanniocalcin (STC) [44]. STC is produced in the corpuscles of Stannius, unique for teleost fish. Besides the uptake of Ca²⁺ in the gills, STC is also involved in the intestinal uptake of Ca²⁺ in freshwater and seawater fish. Hirano [19] reported an increased Ca²⁺ absorption in intestine from freshwater-adapted eel in which the corpuscles of Stannius were surgically removed.

The fish enterocyte provides a unique model with defined endocrine control mechanisms at the apex and basis of the cell. The aim of this study was to analyze the Ca²⁺ transporting mechanism in the apical membrane of the tilapia enterocyte.

Materials and Methods

Sexually mature tilapia (Oreochromis mossambicus) of both sexes, obtained from laboratory stock and weighing 250 to 450 g, were kept in 100-liter aquaria, supplied with running Nijmegen tap water (8-10 mOsmol, [Ca²⁺] = 0.8 mmol · l⁻¹, 25°C) under a photoperiod of 16 hr light : 8 hr darkness. The fish were fed Trouvit® commercial fish food (Trouw, Putten, The Netherlands), 1.5% of the body weight per day.

Materials

⁴CaCl₂ and d-[¹⁴C]mannitol were purchased from Amersham (UK). Trizma-7.0® and Scintillator 299® were from Sigma® (no. T 3503) and Packard, respectively. All other chemicals were analytical grade and obtained from commercial suppliers.
Table. Enrichment and relative recoveries of marker enzyme activities in tilapia intestinal brush border membrane preparations (mean ± SEM)

<table>
<thead>
<tr>
<th>Marker</th>
<th>Enrichment*</th>
<th>Recovery% (±SEM)</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>APase</td>
<td>17.0 ± 3.2</td>
<td>5.1 ± 0.8</td>
<td>9</td>
</tr>
<tr>
<td>(Na⁺ + K⁺)-ATPase</td>
<td>2.8 ± 0.5</td>
<td>0.9 ± 0.2</td>
<td>11</td>
</tr>
<tr>
<td>SDH</td>
<td>0.15 ± 0.07</td>
<td>0.06 ± 0.03</td>
<td>6</td>
</tr>
<tr>
<td>β-D-Glucuronidase</td>
<td>0.7 ± 0.1</td>
<td>0.30 ± 0.04</td>
<td>6</td>
</tr>
<tr>
<td>Protein</td>
<td>0.06 ± 0.03</td>
<td>0.32 ± 0.02</td>
<td>12</td>
</tr>
</tbody>
</table>

* Enrichment is the ratio of the specific activities in the brush border membrane preparation and in the initial homogenate.

** Recovery was calculated as the percentage total activity (equals specific activity × total protein) in the brush border membrane preparation relative to that in the initial homogenate.

† The activity of sucrase, another brush border membrane marker, was not detectable in the initial homogenate. In eight brush border membrane preparations the specific sucrase activity averaged 4.0 ± 0.4 μmol glucose · hr⁻¹ · mg⁻¹ (mean ± SEM).

ISOLATION OF INTESTINAL BRUSH BORDER MEMBRANES

Fish were killed by spinal transection. The peritoneal cavity was opened and the intestinal tract exposed. All subsequent steps were performed at 0–4°C. The proximal 30 cm of the intestine were quickly removed, flushed with ice-cold saline containing 150 mmol · L⁻¹ NaCl, 1 mmol · L⁻¹ N-[2-hydroxyethyl]piperazine-N'-[2-ethanesulphonic acid] (HEPES)/Tris(hydroxy-methyl)aminomethane (Tris), pH 8.0, 1 mmol · L⁻¹ dithiothreitol and 0.1 mmol · L⁻¹ ethylene-diaminetetraacetic acid (EDTA), cut open lengthwise and rinsed with the same saline. The mucosa was scraped off its underlying muscle layers with the aid of a microscope slide. From one fish, around 0.6 g wet weight scrapings were obtained. Scrapings were disrupted by 25 strokes in a glass dounce homogenizer equipped with a loosely fitting pestle in 40 ml homogenization buffer containing 250 mmol · L⁻¹ sucrose, 10 mmol · L⁻¹ HEPES/Tris, pH 8.0, 1 mmol · L⁻¹ dithiothreitol, and 100 U · ml⁻¹ of the proteinase inhibitor aprotinin. This homogenate was centrifuged for 10 min at 1,400 × g. The resulting supernatant (containing about 70% of the total (Na⁺ + K⁺)-ATPase activity present in the initial homogenate) was discarded. The pellet, which contains the brush border membranes, was resuspended in 40 ml homogenization buffer by 25 strokes in a glass dounce homogenizer and centrifuged for 20 min at 1,400 × g as described above. With the resulting supernatant, an additional 20% of the total (Na⁺ + K⁺)-ATPase activity was removed. The pellet was resuspended in 5 ml mannitol buffer containing 300 mmol · L⁻¹ mannitol, 12 mmol · L⁻¹ Tris/HCl, pH 7.1, using a glass dounce homogenizer (3–7 strokes) and passed through cheesecloth. Five volumes of ultrapure water were added to this suspension and the mixture was homogenized by 20 strokes in a glass dounce homogenizer. Solid MgCl₂ was added to obtain a final concentration of 10 mmol · L⁻¹, and the suspension was mildly agitated for 15 min. To collect the brush border membranes, the suspension was centrifuged (3,000 × g, 15 min); the resulting supernatant was spun at 27,000 × g, 30 min. The pellet, containing the brush border membranes, was resuspended by 30 passages through a 23-G needle and collected in 10 ml KCl buffer containing 150 mmol · L⁻¹ KCl, 0.8 mmol · L⁻¹ free Mg²⁺ and 20 mmol · L⁻¹ HEPES/Tris, pH 7.4. Depending on the subsequent assay to be performed, the required amounts of CaCl₂, ethylene glycol-bis-(β-aminoethly ether) (EGTA), N-(2-hydroxyethyl)-ethylenediamine-N',N'-triacetic acid (HEEDTA) and nitrilotriacetic acid (NTA) were added to this KCl buffer, thus allowing loading of the vesicles with the substances mentioned. Finally, the brush border membranes were pelleted by centrifuging at 27,000 × g, 60 min, and resuspended by 30 passages through a 23-G needle in 0.2–0.4 ml assay buffer. The total isolation procedure lasted 4–5 hr. experiments started within 3 hr after isolation.

ENZYME ASSAYS

Protein recovery was 0.32 ± 0.02% (mean ± SEM, n = 12). The membrane preparations contained 0.4–0.8 mg · ml⁻¹ protein and were used on the day of isolation. The marker enzymes used to characterize the membrane preparation were: alkaline phosphatase (APase, EC 3.1.3.1) and succinate (EC 3.2.1.48) for brush border membranes, (Na⁺ + K⁺)-ATPase (EC 3.6.1.3) for basolateral membranes, succinate dehydrogenase (SDH, EC 1.3.99.1) for mitochondria, and β-D-glucuronidase (EC 3.2.1.31) for lysosomes. Assay procedures for APase, (Na⁺ + K⁺)-ATPase and SDH have been described by Flik et al. [12]. The succrase and β-D-glucuronidase activities were assayed according to Dahlqvist [5] and Fishman and Bernfeld [9], respectively. Membrane protein content was determined with a commercial Coomassie Brilliant Blue reagent kit (Biorad), using bovine serum albumin as a reference. Membranes were preincubated with 0.4 mg saponin per mg membrane protein to permeabilize membrane vesicles and to maximize enzyme activity. Data on purification and recovery of marker enzymes are presented in the Table. The brush border membrane preparation is enriched in succrase and 17-fold in APase. Membranes are not contaminated with mitochondria and lysosomes, and only slightly with basolateral membranes. The specific APase activity averaged 339.3 ± 34.9 μmol nitrophenol · hr⁻¹ · mg⁻¹ (mean ± SEM, n = 9), measured at 37°C, pH 10.4. The values for purification and recovery are congruent with those presented by Pelletier et al. [29] who isolated brush border membranes from trout intestine using a calcium precipitation technique. Their values for the specific APase activity, however, are two orders of magnitude lower than those reported here. Titus et al. [42] isolated brush border mem-
branes from tilapia intestine using a comparable procedure. They reported a similar APase enrichment of their preparation as in the present study.

**Intravesicular Space and Membrane Orientation**

Intravesicular space was determined according to Flik et al. [11]. Membrane vesicles (collected in KCl buffer) were equilibrated for 2 hr at 37°C in KCl buffer, containing 0.1 mmol l⁻¹ D-mannitol and labeled with 161 kBq ml⁻¹ D-[¹⁴C]mannitol. The intravesicular space of the brush border membrane vesicles (BBMV), calculated from the equilibrium mannitol content, was 6.0 ± 1.5 µl mg⁻¹ protein (n = 6). The percentage of right-side-out oriented vesicles was determined on the basis of latent glyceraldehyde-3-phosphate dehydrogenase (EC 1.2.1.12) activity using D-glyceraldehyde-3-phosphate as a substrate. The percentage of inside-out oriented vesicles was determined on the basis of latent glyceraldehyde-3-phosphate dehydrogenase activity (EC 3.1.1.7) activity, using acetylthiocholine as a substrate. Procedures were according to Steck and Kant [37]. Unmasking of latent enzyme activity by triton X-100 treatment revealed 99.0 ± 1.3% right-side-out oriented vesicles, and 7.6 ± 5.2% inside-out oriented vesicles (mean values ± SD, n = 4). The high percentage of resealed right-side-out BBMV is consistent with the observations on rat small intestinal BBMV by Haase et al. [17].

**Vesicle Ca²⁺ Transport**

Two approaches were followed to study Ca²⁺ transport across brush border membranes, viz. (i) zero trans uptake and (ii) isotope equilibrium exchange. A rapid filtration technique was used to determine Ca²⁺ content of BBMV. All transport studies were performed in duplicate, blank values were determined in quadruplicate.

**Zero Trans Studies**

In zero trans uptake studies, Ca²⁺ transport is measured in Ca²⁺-free BBMV. Time course analysis and kinetic studies of the initial rate of Ca²⁺ uptake were performed.

Membrane vesicles were collected in Ca²⁺-free KCl buffer. The assay medium consisted of KCl buffer, to which had been added 0.01 to 5 mmol l⁻¹ free Ca²⁺, 0.5 mmol l⁻¹ EGTA, 0.5 mmol l⁻¹ HEEDTA and 0.5 mmol l⁻¹ NTA. The concentration of CaCl₂ (added as CaCl₂) in the assay medium was 0.8–1.0 MBq ml⁻¹. Incubations were carried out at 37°C, membranes and assay medium were prewarmed before incubation. Samples, containing 50 µl of the incubate, were quenched in 1 ml ice-cold stop buffer (containing 150 mmol l⁻¹ KCl, 20 mmol l⁻¹ Trizma-7.0®, pH 7.4, and a concentration La³⁺ at least two times higher than the concentration of free Ca²⁺ in the assay medium). Blanks and samples were prepared as described for the zero trans experiments. The Ca²⁺ content of the membrane vesicles is expressed in dpm per mg membrane protein. Data are corrected by subtracting the blank value from the exchange values.

**Equilibrium Exchange Influx**

After 21 min (more than 8 x 1/2 for isotope exchange influx) 150 µl of the incubate was diluted in 2 ml of (prewarmed) assay buffer, containing 4⁵Ca²⁺ only. When diluted, the vesicles contained approximately 5% of the total radioactivity content in the assay medium. The 4⁵Ca²⁺ concentration in the vesicles was then 30 times higher than in the medium surrounding the vesicles. An isotope exchange in an essentially isotope-free medium is thus measured. Samples (0.4 ml) from this mixture were quenched in 2 ml stop buffer and immediately filtered. Filters were rinsed, dissolved and the radioactivity counted as described above. Experimentally it was not feasible to obtain true blank values, i.e., measurements of the vesicle tracer content at time point zero, for the equilibrium isotope efflux exchange. These data are therefore presented without zero point correction.

**Equilibrium Exchange Efflux**

Membrane vesicles were resuspended in (tracer-free) assay buffer before the last centrifugation step. The same buffer as in the zero trans experiments was used. All subsequent steps of the isolation procedure were then performed in the same assay buffer, thus allowing loading of the vesicles with Ca²⁺. To measure isotope exchange, membrane vesicles were diluted in the same assay buffer to which 0.8–1.0 MBq ml⁻¹ ⁴⁰Ca²⁺ was added. The stop buffer contained 150 mmol l⁻¹ KCl, 20 mmol l⁻¹ Trizma-7.0®, pH 7.4, and a concentration La³⁺ at least two times higher than the concentration of free Ca²⁺ in the assay medium. Time course analysis and kinetic studies of the initial rate of Ca²⁺ uptake were performed.
Thus, the loading procedure was effective, and our assumption of an equilibrium condition with respect to calcium justified. Samples, containing 150 μl from the control incubation, were also quenched in 3 ml ice-cold stop buffer containing 10 mmol • l⁻¹ La³⁺ or 1.0 mmol • l⁻¹ EGTA. Immediately after quenching, as well as two and five minutes thereafter, aliquots of a quenched sample were filtered and analyzed for radioactivity content. No significant decrease of vesicle ⁴⁰Ca²⁺ content with time was found. The use of lanthanum and EGTA as agents to stop Ca²⁺ movements was thus appropriate.

**Calculations and Statistics**

Free Ca²⁺ and Mg²⁺ concentrations were calculated according to Schoenmakers et al. [35], using the computer program CHELATOR which allows correcting metal-chelator stability constants for effects of temperature, ionic strength and proton activity of the medium. The first and second protonations of the chelating compounds were taken into account. Stability constants are from Sillen and Martell [36].

The (initial) Ca²⁺-dependent zero trans Ca²⁺-uptake could be described by a function containing a saturable and a non-saturable term:

\[ J = \frac{J_{\text{max}} \times [S]}{K_m + [S]} + c \times [S] \]  

(1)

where \( [S] \) stands for the free Ca²⁺ concentration in the medium, \( J_{\text{max}} \) represents the limiting flux (i.e., at infinitely high \([S]\)), \( K_m \) is the substrate concentration at which \( J \) is half maximal, and \( c \) is a constant having the dimension of a limiting permeability (i.e., \( J_{\text{max}}/K_m \)). Data from isotope equilibrium exchange influx and efflux experiments were fitted to the exponential equations (2) and (3), respectively:

\[ f(t) = A_1 \cdot (1 - \exp(-k_1 t)) + A_2 \cdot (1 - \exp(-k_2 t)) \]  

(2)

\[ f(t) = A \cdot \exp(-kt) + \text{Offset} \]  

(3)

where \( A \) represents the limit of a component (i.e., ⁴⁰Ca²⁺-counts per mg protein) and \( k \) is a first-order rate constant. The offset represents the tracer content (adsorbed and exchanged tracer) of the BBMV. Subscripts specify parameters of two exponents. Effects of a nonhomogeneity of the BBMV population (i.e., vesicle size, functionally different vesicle subpopulations) on equilibrium exchange influxes and effluxes were analyzed using Hopfer's [21] criteria. Accordingly, \( k_i \), the rate constant for an individual vesicle \( i \), is considered as the product of the rate constant \( r \) of a transporter common in all vesicles, and a "lumped" constant \( h_i \), including the contribution of the area-to-volume-ratio (i.e., vesicle size) of an individual vesicle \( i \) to the actual isotope exchange influx and efflux determined. Then, \( k_i = h_i \cdot r \), and Eqs. (2) and (3) transform to:

\[ f(t) = \sum w_i \cdot \exp(-h_i r t). \]  

(4)

in which \( w_i \) weights the contribution of vesicle \( i \) to the actual macroscopic isotope exchange measured. The transporter's rate constant \( r \) is dependent on experimental variables such as substrate concentration and temperature, whereas the constant \( h_i \) varies only with vesicle size, transporter density and other vesicle specific parameters. Hopfer [21] then showed that, by defining normalized time \( \tau = r \cdot t / t_{50} \) (\( \approx t / t_{1/2} \)), Eq. (4) becomes

\[ f(\tau) = \sum w_i \cdot \exp(-h_i \tau). \]  

(5)

**Results**

Time course studies of zero trans calcium uptake by BBMV are shown in Fig. 1A and B. The uptake of Ca²⁺ in BBMV in the presence of 0.5 mmol • l⁻¹ Ca²⁺ plateaued after 10 min. On the basis of the equilibrium Ca²⁺ content we calculate a vesicular volume of 7.7 ± 1.8 μl • mg⁻¹ protein (mean ± SD),

![Fig. 1. Time course of zero trans Ca²⁺ uptake in BBMV. Means and SEM are shown. (A) Filled circles: 0.5 mmol • l⁻¹ Ca²⁺ (n = 7); open circles: 5.0 mmol • l⁻¹ Ca²⁺ (n = 4). (B) Uptake at 0.5 mmol • l⁻¹ Ca²⁺ during the first minute of the incubation (n = 5).](image)
Fig. 2. Ca\textsuperscript{2+}-dependent Ca\textsuperscript{2+} uptake in BBMV. Uptake was measured during 1-sec incubations. Mean values and SEM for five different membrane preparations are given. The Ca\textsuperscript{2+} concentration in the assay medium was varied from 10 \( \mu \text{mol} \cdot \text{l}^{-1} \) to 5 mmol \( \cdot \text{l}^{-1} \). Data points were fitted to Eq. (1). Estimates of kinetic parameters are: \( J_{\text{max}} = 0.53 \text{nmol Ca}\textsuperscript{2+} \cdot \text{mg}^{-1} \cdot \text{sec}^{-1} \), \( K_m = 5.8 \text{\mu mol} \cdot \text{l}^{-1} \text{Ca}\textsuperscript{2+} \), \( c = 376 \text{name} \cdot \text{sec}^{-1} \cdot \text{mg}^{-1} \). Dotted lines indicate the saturable and nonsaturable component from Eq. (1).

which is not significantly different from the vesicular mannitol space of 6.0 ± 1.5 \( \mu \text{L} \cdot \text{mg}^{-1} \) protein. Obviously the blank value is a proper representation of tracer binding to the vesicle interior and exterior. Zero \( \text{trans} \) Ca\textsuperscript{2+} uptake deviates fast from linearity, and is linear only during the first second of the time course (Fig. 1B). Figure 1A also shows the uptake in the presence of 5.0 mmol \( \cdot \text{l}^{-1} \text{Ca}\textsuperscript{2+} \) in the extravesicular medium. From the two curves describing the time courses of the zero \( \text{trans} \) uptake at two Ca\textsuperscript{2+} concentrations a similar half-time value of 4 min is obtained.

Figure 2 shows the Ca\textsuperscript{2+} dependent zero \( \text{trans} \) Ca\textsuperscript{2+} uptake in BBMV at initial rate. A saturable and a nonsaturable, linear component can be distinguished. The saturable component is described by a Michaelis-Menten equation. The kinetic parameters \( J_{\text{max}} \) and \( K_m \) were calculated to be 0.53 nmol \cdot sec\textsuperscript{-1} \cdot mg\textsuperscript{-1}, and 5.8 \( \text{\mu mol} \cdot \text{l}^{-1} \text{Ca}\textsuperscript{2+} \), respectively. The constant \( c \) in the nonsaturable term \( c \times [S] \) of Eq. (1) is calculated to be 376 \text{name} \cdot \text{sec}^{-1} \cdot \text{mg}^{-1}. From the intersection of the curves describing the saturable and nonsaturable component the \([\text{Ca}\textsuperscript{2+}])\) at which both components contribute equally to the total uptake can be obtained. Figure 2 demonstrates this to be 1.5 mmol\cdot l\textsuperscript{-1}.

Figure 3A and B show the results of equilibrium isotope influxes and effluxes. Data could not be fitted to a single exponential, indicating a heterogeneity of the BBMV population with respect to vesicle size, functional differences of the calcium transporter, or both. Moreover, there is a considerable scattering of the data around a curve of best fit. The half-time values for influx and efflux experiments are estimated to be 0.6 and 2.0 min, respectively. The rate of \( \text{d-[1-}\text{\textsuperscript{14}}\text{C}]\text{mannitol equilibrium exchange (shown in Fig. 4A and B) is slower than the 45Ca}\textsuperscript{2+} exchange rate. Due to the slower rate of exchange, d-[1-\text{\textsuperscript{14}}\text{C}]\text{mannitol movements are more accurately}}
measured and, hence, the rates of exchange influx and efflux are more reliably estimated. Figure 4A and B show the half-time values for D-[1-14C]mannitol exchange influx and efflux to be 2.4 and 2.3 min, respectively. The similarity of these exchange rates validates our assumption of true equilibrium conditions in our experimental setup.

As is shown in Fig. 5A and B, plotting equilibrium exchange data as a function of normalized time \( \tau \) reduces the scattering. Data points are now conveniently described by a single curve. This indicates that the BBMV population consists of "similar" [21] vesicles, i.e., vesicles differing in size or transporter density only. More importantly, the process of equilibrium exchange, measured over a range of 0.1 to 10 mmol \( \cdot \) l\(^{-1} \) Ca\(^{2+} \), does not show saturation. The lack of a correlation between the exchange rate and the substrate concentration is indicative of a
passive diffusion process or for the activity of a transporter operating well below its level of saturation.

The rate constant of exchange is directly proportional to the surface area of a vesicle and the permeability of a membrane, and inversely proportional to the vesicle’s volume. A check for passive diffusion of a substrate can then be performed by measuring substrate uptake in vesicles with different area-to-volume ratios. Figure 6 shows equilibrium influxes in BBMV, measured at increasing osmolarities of the extravesicular medium. Influx curves are superimposable for all conditions employed. Any change in vesicular volume would have been indicated by different exchange rates and different limits of the curves describing these tracer exchanges. The BBMV do not behave as osmometers, i.e., they do not shrink predictably as a response to an increase in external osmolarity.

**Discussion**

The Ca\(^{2+}\)-dependent zero trans uptake in tilapia intestinal BBMV is a curvilinear process. Similar kinetics for Ca\(^{2+}\) uptake were observed in BBMV of hamster duodenum and ileum [23, 32], rat small intestine [14, 18, 26, 33, 45], chick duodenum [31] and human small intestine [15]. In general curvilinear patterns for the uptake of electrolytes as well as nonelectrolytes are found. This was shown to be the case for e.g., the uptake of Zn\(^{2+}\) in BBMV from pig small intestine [40], and the uptake of acetate, D-glucose and L-proline in tilapia BBMV [42]. Since a curvilinear uptake is displayed by many hydrophilic solutes [3], this does not seem to be a specific property of brush border membranes.

In stripped intestinal mucosa from tilapia, and in the presence of 1.25 mmol L\(^{-1}\) Ca\(^{2+}\) on the mucosal side of the preparation, a net transepithelial Ca\(^{2+}\) flux of 34 ± 8 nmol hr\(^{-1}\) cm\(^{-2}\) was measured [11]. From the intravesicular volume of 7.7 µl mg\(^{-1}\) for BBMV, a vesicle membrane area of 18.9 µm\(^2\) mg\(^{-1}\) is calculated. Assuming that the luminal area represents brush border membrane, 1 cm\(^2\) of intestinal lumen is then equivalent to 529 mg brush border membrane protein. The net transepithelial flux, as measured by Flik et al. [11], can then be expressed as 5 nmol Ca\(^{2+}\) sec\(^{-1}\) mg\(^{-1}\). Figure 2 shows the uptake in BBMV at 1.25 mmol L\(^{-1}\) Ca\(^{2+}\) to be 1 nmol Ca\(^{2+}\) sec\(^{-1}\) mg\(^{-1}\), five times lower than the net transepithelial flux. Such a difference can be anticipated, considering the folding of the lumen and the presence of intestinal microvilli in tilapia.

Zero trans uptakes can be biased by binding of Ca\(^{2+}\) to charged membrane surface areas [21, 43]. This can result in a virtually Ca\(^{2+}\)-deplete intravesicular medium, and hence an artificial prolonged linear phase in the transmembrane movement of Ca\(^{2+}\). From the uptake of Ca\(^{2+}\) and mannitol into BBMV we concluded that Ca\(^{2+}\) binding to the vesicle interior and exterior was appropriately corrected for. Therefore, our measurements seem to represent true initial uptake rates. The curvilinearity of the Ca\(^{2+}\)-dependent uptake in BBMV is usually explained by the presence of a saturable (or channel- or carrier-mediated) component and a nonsaturable (or diffusional) component. Many investigators have focused on the saturable component only, which is considered to be the regulated mechanism in brush border membrane Ca\(^{2+}\) entry. We obtained a Ca\(^{2+}\) concentration of 1.5 mmol L\(^{-1}\) at which the contribution of each component to the total Ca\(^{2+}\) uptake is equal. Luminal Ca\(^{2+}\) concentrations are in the range of 5–25 mmol L\(^{-1}\) (personal observation). The saturable component is then operating under \(J_{\text{max}}\) conditions, and, having a \(K_m\) in the micromolar range, is obviously not the rate-limiting step in brush border membrane transport. The question then arises whether the nonsaturable component is rate limiting and whether this is a true diffusion process. Indeed, Christensen [3] suggested that an apparently linear portion of a curve describing uptake rate as a function of substrate concentration could well be a part of a second rec-

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**Fig. 6.** Effect of external osmolarity on \(^{45}\)Ca\(^{2+}\) equilibrium influx in BBMV. During the final isolation steps, BBMV were homogenized and collected in 100 mOsmol lactose and 5.0 mmol L\(^{-1}\) Ca\(^{2+}\). Membrane vesicles were incubated in uptake media containing sufficient lactose to reach the indicated osmolarity. Other assay conditions are the same as for equilibrium exchange, as described in Materials and Methods. (○ 152 mOsmol, □ 253 mOsmol, △ 286 mOsmol).
tangular hyperbola characterizing a second saturable component.

Information on the nonsaturable component is contained in the constant $c$ of Eq. (1), for tilapia BBMV calculated to be 376 nl · sec$^{-1}$ · mg$^{-1}$. This constant $c$ has the same dimension as the ratio $J/[S]$, i.e., initial rate flux divided by the substrate concentration, which is equivalent to the definition of the permeability coefficient $P_c$, where $P = J/(C_i - C_o)^{-1}$. Here, $J$ is the net flux, $C_i$ and $C_o$ are the substrate concentrations on the two sides of the membrane, and $A$ is the membrane area. Since $c = P · A$, dividing $c$ by the vesicle membrane area gives a Ca$^{2+}$ permeability coefficient, $P_c$, of $2 \cdot 10^{-3}$ cm · sec$^{-1}$. Gunther et al. [16] reported an average permeability of 5 nl · mg$^{-1}$ · sec$^{-1}$ and a permeability coefficient of approximately $10^{-5}$ cm · sec$^{-1}$ for a number of monovalent cations in rabbit jejunal BBMV. Compared with basal permeabilities (i.e., the permeability in the absence of a specific transport pathway) for a number of nonelectrolytes, as listed by Stein [38], the calculated Ca$^{2+}$ permeability is very high, and is not compatible with a simple diffusion of Ca$^{2+}$. The identical uptake rates as determined at 0.5 and 5.0 mmol · l$^{-1}$ Ca$^{2+}$ also do not agree with diffusion of Ca$^{2+}$ through a lipid bilayer. Diffusion of Ca$^{2+}$ through a lipid bilayer is energetically unfavorable, due to the hydrophilic nature of the Ca$^{2+}$ ion and, hence, a high degree of hydration. Permeability coefficients of synthetic lipid bilayers for charged molecules are in the range of $10^{-10}$–$10^{-12}$ cm · sec$^{-1}$. A protein-mediated pathway can explain the high Ca$^{2+}$ permeability of the tilapia intestinal brush border membrane. Moreover, the presence of a transporter may form a mechanism to control the steep Ca$^{2+}$ gradient across the brush border membrane, and for its endocrine control.

The nonsaturable component was further analyzed using isotope equilibrium exchange. During the isolation and loading procedure, membrane vesicles were equilibrated in an essentially infinitely large pool of $^{40}$Ca$^{2+}$, thus allowing masking of the membrane binding sites. The first-order rate constant describing the equilibrium exchange, $k$, is equal to $J_{\text{ex}0}/([K_m + [S]])$ [7]. We found $k$ to be independent of [S], indicating $K_m \gg [S]$ for Ca$^{2+}$ concentrations up to 10 mmol · l$^{-1}$, so that $k = J_{\text{ex}0}/K_m$. The nonsaturable component appears not to be a procedural artifact, but a property of the Ca$^{2+}$ movement across the tilapia intestinal brush border membrane.

The results presented so far lead us to postulate a low-affinity Ca$^{2+}$ transporting mechanism with an allosteric regulatory Ca$^{2+}$ binding site. The mechanism of Ca$^{2+}$ entry across brush border membranes in general is still unknown. The existence of Ca$^{2+}$ channels in brush border membranes has not yet been documented [2]. Wilson et al. [45] found an acceleration of Ca$^{2+}$ transport in rat intestinal BBMV by strontium, and proposed a carrier mechanism for the transport of Ca$^{2+}$. Based on the observation of many vacuoles in small intestinal enterocytes, Pansu et al. [27] suggested the nonsaturable Ca$^{2+}$ transport in young rats to be pinocytosis-like. However, such a shuttle mechanism for transcellular Ca$^{2+}$ transport cannot, in our opinion, explain the Ca$^{2+}$ permeability found by the same authors in older rats lacking vacuoles in their enterocytes, nor the high Ca$^{2+}$ permeability found in isolated intestinal brush border membranes from tilapia and other animal species.

The results of the equilibrium experiments also permit the conclusion that the tilapia intestine shows no regional differences along the longitudinal axis and the mucosa-serosa axis. Functionally different vesicle subpopulations, e.g., derived from intestinal regions having Ca$^{2+}$ transporters with different kinetic characteristics, would have become visible by nonsuperimposable curves. The homogeneity of the tilapia intestine is in contrast to rat and hamster data, where Ca$^{2+}$ uptake characteristics were different for BBMV preparations from proximal and distal intestinal regions [23, 28, 32]. Stieger and Murer [39] isolated BBMV from rat small intestine by a Mg/EGTA precipitation method, and performed an additional fractionation by free-flow electrophoresis and sucrose density gradient centrifugation. They found different subfractions of BBMV in the whole intestine, as well as in duodenum, ileum and jejunum with respect to enzyme properties, and are probably under hormonal control.

Tilapia BBMV did not adjust their volumes to a change in external osmolarity, indicative for a water tightness of the intestinal brush border membrane. In euryhaline fish such as the Japanese eel (Anguilla japonica) kept in freshwater a high prolactin secretion is correlated with a decreased net water influx, indicating the intestine as a prolactin target [20]. Dave [6] reviews evidence for a prostaglandin-mediated pathway by which prolactin modifies membrane lipid microviscosity. Tilapia enterocytes could well be a target for serum prolactin, explaining the impermeability of BBMV to water.
The authors wish to thank Tom Spanings for his superb organization of fish husbandry, and Maarten de Jong (Dept. of Physiology, Faculty of Medicine, University of Nijmegen) for making the automated stopped-flow apparatus available to us.

References


Received 10 July 1992; revised 2 October 1992