Acute ENaC Stimulation by cAMP in a Kidney Cell Line is Mediated by Exocytic Insertion from a Recycling Channel Pool

MICHAEL B. BUTTERWORTH,1 ROBERT S. EDINGER,2 JOHN P. JOHNSON,2 and RAYMOND A. FRIZZELL1

1Department of Cell Biology and Physiology and 2Department of Medicine, Renal-Electrolyte Division, School of Medicine, University of Pittsburgh, Pittsburgh, PA 15261

ABSTRACT Acute hormonal regulation of the epithelial sodium channel (ENaC) in tight epithelia increases tranacellular Na+ transport via trafficking of intracellular channels to the apical surface. The fate of the channels removed from the apical surface following agonist washout is less clear. By repetitively stimulating polarized mouse cortical collecting duct (mCCD, mPCCD14) epithelia, we evaluated the hypothesis that ENaC recycles through an intracellular pool to be available for reinsertion into the apical membrane. Short circuit current (Isc), membrane capacitance (Cm), and conductance (Gf) were recorded from mCCD epithelia mounted in modified Ussing chambers. Surface biotinylation of ENaC demonstrated an increase in channel number in the apical membrane following cAMP stimulation. This increase was accompanied by a 83 ± 6% (n = 31) increase in Isc and a 15.3 ± 1.5% (n = 15) increase in Gf. Selective membrane permeabilization demonstrated that the Cm increase was due to an increase in apical membrane capacitance. Isc and Cm declined to basal levels on stimulus washout. Repetitive cAMP stimulation and washout (~1 h each cycle) resulted in response fatigue; ΔIsc decreased ~10% per stimulation–recovery cycle. When channel production was blocked by cycloheximide, ΔIsc decreased ~15% per stimulation cycle, indicating that newly synthesized ENaC contributed a relatively small fraction of the channels mobilized to the apical membrane. Selective block of surface ENaC by benzamil demonstrated that channels inserted from a subapical pool made up >90% of the stimulated Isc, and that on restimulation a large proportion of channels retrieved from the apical surface were reinserted into the apical membrane. Channel recycling was disrupted by brefeldin A, which inhibited ENaC exocytosis, by chloroquine, which inhibited ENaC endocytosis and recycling, and by latrunculin A, which blocked ENaC exocytosis. A compartment model featuring channel populations in the apical membrane and intracellular recycling pool provided an adequate kinetic description of the Isc responses to repetitive stimulation. The model supports the concept of ENaC recycling in response to repetitive CAMP stimulation.

KEY WORDS: cellular traffic • recycle • short-circuit current • capacitance • biotinylation

INTRODUCTION

The epithelial sodium channel (ENaC) is localized at the apical membranes of several epithelia, including lung, colon, sweat gland, salivary glands, taste buds, principal cells of the cortical collecting duct (CCD), and the collecting tubules of mammalian nephrons (Garty and Palmer, 1997; Rossier et al., 2002). In the renal CCD, ENaC constitutes the regulated, rate-limiting entry step in Na+ reabsorption, and its activity is under strict hormonal control. Abnormalities in ENaC regulation and activity have been linked to a number of human disorders, including hypertension (Liddle’s disease), pseudo-hypo aldosteronism (PHA), and cystic fibrosis (Liddle et al., 1963; Hanukoglu, 1991; Mall et al., 2004). The channel is comprised of three homologous subunits: α, β, and γ, whose stoichiometry is thought to conform to the ratio 2α:1β:1γ (Firsov et al., 1996, 1998; Gormley et al., 2003). ENaC exhibits characteristic biophysical properties, including a low single channel conductance (~5 pS), relatively long open and closed times, a high Na+ to K+ selectivity ratio (>100:1), and is blocked by the pyrazine diuretic, amiloride, at submicromolar concentrations (Kellenberger and Schild, 2002). The expression and activity of ENaC is regulated by numerous factors, including long (aldosterone) and short (vasopressin, insulin) term hormonal regulation, and sensitivities to flow, stretch, proteases, and altered intra- and extracellular Na+ concentrations (Palmer, 1992; Garty and Palmer, 1997).

In many ENaC-expressing tissues and model epithelia, a rapid stimulation of sodium transport is elicited by vasopressin through increases in intracellular cAMP levels and PKA activation (Benos et al., 1995; Garty and Palmer, 1997; Els and Butterworth, 1998; Ecelbarger et al., 2001; Snyder, 2002). Some debate exists regarding the mechanism by which vasopressin increases Na+ transport (Rossier, 2002); that is, increased open

*Correspondence to Michael B. Butterworth: michael7@pitt.edu

Abbreviations used in this paper: BFA, brefeldin A; CCD, cortical collecting duct; CHX, cycloheximide; ENaC, epithelial sodium channel; LatA, latrunculin A; TGN, trans-Golgi network.
probability ($P_o$) or increased apical membrane channel number (N). Nevertheless, several studies have suggested that the acute (sec–min) increase in Na$^+$ transport is the result of regulated insertion of ENaC-containing intracellular vesicles into the apical membrane (increased N) (Erlilj et al., 1999; Snyder, 2000; Weisz et al., 2000; Butterworth et al., 2001; Morris and Schafer, 2002; Planes et al., 2002). The residence time of channels at the apical surface is regulated by removal and degradation processes, which are initiated by the binding of Nedd4-2 to the cytoplasmic COOH termini of the ENaC subunits. All three subunits contain a proline rich region (PPXXYXXL), which binds Nedd4 to target ENaC for ubiquitin-dependent internalization and degradation (Goulet et al., 1998; Abriel et al., 1999; Farr et al., 2000; Staub et al., 2000; Malik et al., 2001). Mutations in the PY motifs at the subunit COOH termini of β- and γ-ENaC result in increased ENaC residency in the plasma membrane and a constitutive increase in Na$^+$ transport. Although polyubiquitination is often a marker for proteosomal degradation, monoubiquitination is also associated with endocytosis and recycling of plasma membrane proteins such as the epidermal growth factor (EGF) receptor (Sorkin, 2001; Dikic, 2003). Thus, the extent to which ENaC retrieved from the apical surface is subjected to degradation versus recycling is unclear.

To characterize the acute regulation of ENaC by cAMP, mouse CCD cells cultured on permeable supports were repeatedly stimulated to fatigue the short-circuit current ($I_{sc}$) response and identify the mechanisms regulating apical ENaC activity. We show that PKA activation resulted in the insertion of ENaC subunits into the apical surface, which results in cAMP-induced increases in $I_{sc}$ and epithelial capacitance ($C_T$). On agonist removal, both $I_{sc}$ and $C_T$ declined to basal levels, and subsequently the cells were able to respond to additional rounds of stimulation. By blocking protein biosynthetic pathway or membrane trafficking pathways, evidence was provided for a subapical pool of recycling channels. We propose that distinct ENaC populations in CCD epithelia account for constitutive and regulated Na$^+$ transport and that the acute stimulation of Na$^+$ transport is due primarily to insertion of ENaC from a tightly regulated channel recycling pool.

MATERIALS AND METHODS

Cell Culture

The mpkCCD14 cells (provided by A. Vandewalle and M. Bens, Institut National de la Santé et de la Recherche Médicale, Paris, France) were grown in flasks (passage 30–40) in defined medium as described previously (Vinciguerra et al., 2003). Growth medium was composed of equal volumes DMEM and Ham’s F12, 60 mM sodium selenate, 5 μg/ml transferrin, 2 mM glutamine, 50 nM dexamethasone, 1 nM triiodothyronine, 10 ng/ml epidermal growth factor, 5 μg/ml insulin, 20 mM d-glucose, 2% vol/vol FCS, and 20 mM HEPES (Invitrogen, GIBCO BRL, Sigma-Aldrich), pH 7.4, at 37°C in 5% CO$_2$/95% air atmosphere. The medium was changed every second day. For all electrophysiological experiments, the CCD cells were subcultured onto permeable filter supports (0.4 μm pore size, 0.33 cm$^2$ surface area; Transwell, Corning Costar). Cells were kept on filters for at least 7 d in defined medium which was changed every second day. Typically after 7 d, a confluent transporting cell monolayer had developed that could be assessed by recording open circuit voltage and trans epithelial resistance using “chopstick” electrodes (Millipore). At least 24 h before use in any investigation, medium incubating cells on filters was replaced with a minimal medium (without drugs or hormones) that contained only DMEM and Ham’s F12.

Standard $I_{sc}$ Current Recordings

Cells cultured on filter supports were mounted in modified Costar Ussing chambers, and the cultures were continuously short circuited with an automatic voltage clamp (Department of Bioengineering, University of Iowa, Iowa City, IA). Transepithelial resistance was measured by periodically applying a 2.5-nA bipolar pulse and calculated using Ohm’s law. The bathing solutions were composed of 120 mM NaCl, 25 mM NaHCO$_3$, 3.3 mM KH$_2$PO$_4$, 0.8 mM K$_2$HPO$_4$, 1.2 mM MgCl$_2$, 1.2 mM CaCl$_2$, and 10 mM glucose. Chambers were constantly gassed with a mixture of 95% O$_2$/5% CO$_2$ at 37°C, which maintained the pH at 7.4. To facilitate chamber washes, the bases of both chamber compartments were connected to a peristaltic pump, in order to control volume flow rate, with fluid removed from the top of the chamber. At the minimum, a fivefold volume exchange (25–30 ml) was performed to wash out the chamber, and in some cases, this was increased (see below) to ensure sufficient removal of agonist or other compounds. As the cells were sensitive to changes in pressure, flow, and temperature, the chamber solutions were only exchanged during wash periods with no flow at steady states. A disruption was always noted in the recorded traces during basolateral chamber wash periods, but currents returned to prewash levels at the end of the fluid exchange. Similar transients were observed when solution of the same composition was exchanged in the basolateral chamber. A typical stimulation cycle lasted 1 h and involved the addition of 10 μM forskolin basolaterally, which produced a maximum $I_{sc}$ stimulation after 20–30 min; forskolin was washed from the basolateral side of the chamber and current declined back to basal levels within 30 min. To determine the net Na$^+$ transport through ENaC, 10 μM amiloride was added to the apical cell surface at the end of each experiment.

Simultaneous $I_{sc}$, Capacitance, and Impedance Recordings

The same modified Ussing chambers were used to record simultaneous $I_{sc}$ and $C_T$ traces. The voltage clamp and recording system used to acquire data were designed and constructed by W. Van Driessche (Katholieke Universiteit, Leuven, Belgium) and have been described in detail previously (Erlilj et al., 1999; Weber et al., 1999). In brief, hardware for $C_T$ measurements used two Digital Signal Processing (DSP) boards (Model 310B; Dalanco Spry) with one board used to record transepithelial conductance ($G_T$) and short-circuit current ($I_{sc}$) and the second, $C_T$. Due to the requirement to generate multiple sine waves in order to obtain simultaneous measurements, the system only updated recorded values every 7 s. As stimulation events occurred over tens of minutes, this was not limiting. Continuous $C_T$ values were calculated from imposed voltage sine waves of frequencies 2, 2.7, 4.1, 5.4, and 8.2 kHz. The 4.1-kHz trace is presented in all $C_T$ figures. Impedance analysis was performed at steady states by simultaneously imposing 78 sine waves to the command input of the...
voltage clamp and analyzed as previously described (Weber et al., 1999). The Nyquist plot obtained from the impedance spectra exhibited only one semicircle, which is presumed to reflect the apical membrane capacitance. The relative contribution of each membrane to \( C_T \) was determined by selectively permeabilizing each membrane with ionophores (see below). It was not practical to utilize ionophores in long term studies performed here, and as it could be demonstrated by permeabilization studies that the capacitance changes were the result of alterations in \( C_A \) alone (see below), \( C_B \) recordings could be assumed to reflect changes in \( C_A \) (see also Erli et al., 1999; Weber et al., 1999).

**Membrane Permeabilization**

To isolate apical and basolateral membranes electrically, bathing ringers solutions were kept identical on both sides of the chamber and increasing doses of nystatin (25, 50, 75, 100 \( \mu \)M; Sigma-Aldrich) were added to either side to incrementally permeabilize the membrane and assess the effect by monitoring \( I_{sc} \) and \( C_T \). To investigate the effect of forskolin stimulation on capacitance changes in either membrane, 100 \( \mu \)M nystatin was used, as full membrane permeabilization was achieved at this concentration.

**Surface Biotinylation**

CCD cells cultured on 75-mm diameter filter supports (Transwell) were washed (5 min) with ice-cold PBS with agitation on ice to remove growth media. The apical membrane was biotinylated using 0.5 mg/ml Streptavidin (Pierce Chemical Co.) in borate buffer (85 mM NaCl, 4 mM KCl, 15 mM Na\(_2\)HPO\(_4\), pH 9) for 20 min. The basolateral surface was incubated in growth medium containing FBS to prevent biotinylation. Biotinylation was quenched by adding a double volume FBS-containing medium. Samples were collected in 2 ml of protein concentration of the post nuclear supernatant was determined, and 200 \( \mu \)g of protein was combined with a streptavidin bead slurry (Pierce Chemical Co.) and incubated overnight at 4°C. Samples from the streptavidin beads were collected in 2% sample buffer containing 10% \( \beta \)-mercaptoethanol and incubated for 20 min at room temperature. Samples were heated at 95°C for 3 min, separated by SDS-PAGE, and subjected to Western blot analysis with the appropriate antibodies and visualized using chemiluminescence (Perkin-Elmer). To verify that intracellular proteins were not biotinylated during the experimental procedure, biotinylated samples collected from unstimulated cells were processed using Metamorph (Universal Imaging Corp.) with adjustments made to only contrast and brightness (phalloidin staining) or simple nearest neighbor deconvolution followed by brightness adjustment (GFP-Endo).

**Microscopic Image Capture and Processing**

All fluorescent images were acquired on a Nikon Diaphot-300 epifluorescent microscope using a 40X 1.4 A.A. objective and equipped with a Hamamatsu C4742-95 CCD camera. Images were processed using Metamorph (Universal Imaging Corp.) with adjustments made to only contrast and brightness (phalloidin staining) or simple nearest neighbor deconvolution followed by brightness adjustment (GFP-Endo).

**Reagents, Fluorophores, and Antibodies**

Brefeldin A (BFA) and Latrunculin A (LatA), which were reconstituted in DMSO at 1,000 times stock solutions, and Alexa Fluor 568 phalloidin were obtained from Molecular Probes. CHX (500 mg/ml stock solution in DMSO), arginine-vasopressin (100 IU/ml stock), benzamil, nickel chloride, chloroquine, and Hoechst nuclear stain were obtained from Sigma-Aldrich. Forskolin (10 mM stock in ethanol) was obtained from Calbiochem (CN Biosciences). DMSO and ethanol, used as vehicles, had no significant effect on recorded \( I_{sc} \) when added at 1:1,000 dilution alone (not depicted). ENaC antibodies were donated by K. Peters (Department of Cell Biology, University of Pittsburgh; \( \alpha \)-ENaC) and M. Knepper (Laboratory of Kidney and Electrolyte Metabolism, National Heart, Lung, and Blood Institute, National Institutes of Health, Bethesda, MD; \( \beta \)-ENaC) or were commercially obtained for \( \gamma \)-ENaC (Abcam).

**Data Analysis and Modeling Software**

Summarized data are presented as mean and standard error (Sigmaplot 2000; SPSS). Simple linear regression fits were performed using Sigmaplot. Object-based modeling software (Model Maker V4; Model Kinetix) was used to construct a compartmental model to describe channel trafficking events. Traces for the kinetic events were obtained by using the built-in integration engines provided in the software and integrating the functions over desired time lengths.

**RESULTS**

**cAMP Stimulation Increases \( I_{sc} \) and \( C_T \)**

Fig. 1 illustrates typical short-circuit current (\( I_{sc} \)) and transepithelial capacitance (\( C_T \)) traces from CCD epithelia stimulated with 10 \( \mu \)M forskolin. Basal \( I_{sc} \) increased, after an initial dip due to \( K^+ \) efflux (the small \( I_{sc} \) decline was eliminated in the presence of 5 mM Met and Cys in the same media for 15 min before addition of 50–100 \( \mu \)g of [\(^{35}\)S]Met/Cys (New England Nuclear, PerkinElmer) and pulsed for 15 min. CHX-treated cells were treated the same way except that media was supplemented with 200 \( \mu \)g/ml CHX. The cells were washed and isolated as described above. Equal amounts of lysate were separated by SDS-PAGE. Total radioisotope incorporation was quantitated from the dried gel using a phosphoimager (Bio-Rad Laboratories).
BaCl₂; unpublished data), to a value almost twofold greater than baseline over a 25-min time course. The increase in I\textsubscript{sc} was immediately preceded, and then paralleled, by an increase in C\textsubscript{T}. Addition of 10 μM amiloride reduced the I\textsubscript{sc} by ~90%, demonstrating that the majority of the transepithelial current was ENaC-mediated INa. Amiloride had no significant effect on C\textsubscript{T}, indicating that the measured capacitance was not influenced by changes in transepithelial current or conductance. A similar response was observed when epithelia were stimulated with arginine vasopressin (AVP), however a larger stimulation, which was more readily reversed, was recorded with forskolin (unpublished data). As a result, all subsequent experiments made use of forskolin to elicit cAMP-mediated increases in I\textsubscript{Na}.

**Forskolin Increases Apical Membrane ENaC Density**

To investigate the basis of the increased I\textsubscript{Na}, apical biotinylation was performed on filter-cultured CCD epithelia using identical stimulation conditions. Increases in all ENaC subunits accessible to biotin labeling at the apical surface were observed following a 30-min forskolin stimulation (Fig. 2 A, with densitometric quantitation of forskolin-induced fold increases in signal presented beneath the blot for each subunit). This indicated that the increase in I\textsubscript{sc} was due, in part, to insertion of channel subunits that increased apical ENaC density. As a control, no intracellular biotinylation was observed when samples were probed for actin (unpublished data). As a result, all subsequent experiments made use of forskolin to elicit cAMP-mediated increases in I\textsubscript{Na}.

**Forskolin Increases Apical Membrane Capacitance**

To better interpret the C\textsubscript{T} measurements, the epithelium was modeled as series RC circuits corresponding to the apical and basolateral membranes, shunted by a paracellular resistance. In this standard lumped model, C\textsubscript{T} is described by

\[
\frac{1}{C_T} = \frac{1}{C_A} + \frac{1}{C_B}
\]

(Weber et al., 1999; Erlij et al., 1999).

In polarized A6 epithelia, C\textsubscript{B} is at least six times larger than C\textsubscript{A} so that the relative contribution of C\textsubscript{B} to transepithelial capacitance is small (Erlij et al., 1994, 1999; Wills et al., 1992). Accordingly, changes in C\textsubscript{T} primarily reflect changes in C\textsubscript{A} (Weber et al., 1999; Pausnecu and Helman, 2001a,b).

The individual membrane contributions to the forskolin-induced ΔC\textsubscript{T} across CCD epithelia were evalu-
ated using selective membrane permeabilization with the addition of 25–100 μM nystatin to either the apical or basolateral bath. Incremental apical nystatin additions were used to resolve the basolateral membrane electrically while simultaneously monitoring \( I_{SC} \), conductance, impedance, and capacitance to determine the point at which the apical surface was effectively permeabilized. The dose-dependent progression to an isolated basolateral membrane is demonstrated by the impedance plots shown in Fig. 3 A. Apical permeabilization as illustrated for a typical experiment in Fig. 3 B resulted in (a) effective elimination of the apical membrane resistance, reflected by a decrease in \( R_T \) from \( \sim 3,000 \ \Omega \cdot \text{cm}^2 \) to \( \sim 400 \ \Omega \cdot \text{cm}^2 \), (b) an increase in \( I_{SC} \), which was blocked by ouabain (not depicted), to a value six to sevenfold greater than the basal current, this ouabain-sensitive current is due to cation transport of the basolateral \( \text{Na}^+ / \text{K}^+ \text{ATPase} \) and has been previously demonstrated for polarized epithelial cells (Fujii and Katz, 1989; Rokaw et al., 1996; Ito et al., 1999), and (c) an increase in \( C_T \) to a value about fourfold higher than that observed for the intact epithelium. The \( C_T \) value following apical nystatin treatment verifies the assumption that the basal surface area was far larger than that of the apical membrane, so that in the intact epithelium, \( C_A \) dominates the \( C_T \) measurement. Four similar treatments produced nystatin-induced increases in \( C_T \) that averaged 4.15 ± 0.12-fold. Substituting the capacitance values observed in Fig. 3 B into Eq. 1 (above), a calculated \( C_A = 1.25 \ \mu \text{F/cm}^2 \) is obtained, which is similar to \( C_T \) values of 1.29 ± 0.03 \( \mu \text{F/cm}^2 \) recorded in control, unstimulated epithelia (\( n = 15 \)). Moreover, in the presence of apical nystatin, there was no significant change in recorded capacitance in response to forskolin, suggesting that a basolateral capacitance change did not contribute significantly to the forskolin-induced increase in \( C_T \). Conversely, when the basolateral membrane was permeabilized, apical capacitance increased 14 ± 2% \( (n = 4) \), which was not significantly different from the \( C_T \) increase of 15 ± 1.5% \( (n = 15) \) observed in control, unpermeabilized epithelia (Fig. 3 C).

Repetitive cAMP Stimulation Causes \( I_{SC} \) Response Rundown

The \( I_{SC} \) and \( C_T \) responses to repetitive forskolin challenges were examined in an attempt to determine whether this technique could be used to evaluate the fate of inserted channels following washout of the cAMP agonist. Typical \( I_{SC} \) and \( C_T \) traces from one such experiment are presented in Fig. 4 A. With each successive stimulation–recovery cycle (\( \sim 1 \) h each), the forskolin-induced \( \Delta I_{SC} \) diminished. The resulting \( \Delta I_{SC} \) decay is plotted in Fig. 4 B, where the slope of the linear regression fit indicates a loss of \( \sim 10% \) of the \( \Delta I_{SC} \) with each response (slope = -11.33, \( r^2 = 0.99 \)). Plotted on

![Figure 3. Membrane permeabilization. (A) Nyquist plots from a CCD epithelium to which increasing concentrations of apical nystatin was added. From an unpermeabilized state, a single impedance locus (one semicircle) transitioned through two loci until only one locus corresponding to the basolateral membrane was observed (at 100 μM nystatin). (B) Time course of \( I_{SC} \) (black) and \( C_T \) (gray) plots in response to apical nystatin addition (white bar). Small changes in recorded basolateral capacitance with forskolin (gray bar) were insufficient to account for observed \( \Delta C_T \) changes. (C) A similar plot (as in B) for basolateral permeabilization exhibits a \( C_T \) response to 10 μM basolateral forskolin addition similar to unpermeabilized epithelia. \( I_{SC} \) decreased due to removal of transepithelial driving force. Small drift in \( G_T \) (open circles) indicates that transepithelial resistance remained fairly constant.](image-url)
Acute ENaC Regulation by Recycling from an Intracellular Pool

separate axes, the change in forskolin-induced $C_T$ also fell (slope = $-1.47$, $r^2 = 0.95$). By plotting $\%\Delta ISC$ vs. $\%\Delta C_T$ for each stimulation cycle (Fig. 4 B, inset), it is clear from the single regression analysis that the decline in $\Delta ISC$ was correlated with the decline in $\Delta C_T$.

Retrieved ENaC Recycles to the Apical Membrane on Restimulation

From these data, it is apparent that these cells respond repeatedly to forskolin, which elicits an insertion of channels into the apical membrane. To determine whether these responses involve insertion of recycled channels, ENaC-dependent $ISC$ was blocked in two ways. Partial, irreversible inhibition of ENaC-Na$^+$ currents expressed in oocytes by NiCl$_2$ has been demonstrated previously (Sheng et al., 2002). Addition of 20 mM NiCl$_2$ apically resulted in a significant $26.1 \pm 4.5\%$ ($n = 5$) inhibition in forskolin-stimulated $ISC$ (Fig. 5 A), which could not be reversed by apical washing (not depicted). Restimulation of epithelia following a 30-min recovery period produced a significantly reduced $ISC$ response to forskolin compared with uninhibited controls, suggesting that channels that were inserted into the membrane were those previously inhibited by NiCl$_2$ (Fig. 5 B).

Similarly, 10 µM benzamil was used to selectively block channels that reside in the apical membrane. Benzamil is more potent than amiloride, and it is not readily reversible, due to a much slower dissociation rate relative to that of amiloride (Kleyman et al., 1986; Kellenberger et al., 2003) and the fact that it was not washed out of the apical membranes of mCCD epithelia mounted in the Ussing chambers (compare with amiloride washout; see Fig. 6, D–F). Benzamil was added at different times during the stimulation protocol to block membrane-res-
ident ENaC. Examples of $I_{SC}$ traces from two such experiments are presented in Fig. 6 (A and B). As a control, the cells were initially subjected to one round of forskolin stimulation. In Fig. 6 A, benzamil was added at baseline following forskolin washout, while in Fig. 6 B, it was added at the peak of the next forskolin response. After 5 min incubation with benzamil, the apical chamber was washed extensively (10-fold volume exchange as opposed to the usual fivefold) to completely remove the blocker from the bath. In Fig. 6 A, epithelia were then restimulated with forskolin after benzamil had blocked the basal current. The observed increase in $I_{SC}$ was therefore due to insertion of channels from an intracellular pool that was inaccessible to apical benzamil blockade. The increase in $I_{SC}$ supports the biotinylation data (Fig. 2), which detected additional ENaC subunits at the apical surface following forskolin stimulation. In Fig. 6 B, channels were blocked by benzamil addition at the peak of the forskolin response. After forskolin washout, and allowing sufficient time to ensure retrieval of channels from the apical surface (30 min; see Fig. 4 A), the cells were restimulated. The $I_{SC}$ response to forskolin was essentially abolished, suggesting that if channels were recruited to the apical surface, they were those that had been blocked by benzamil during the previous stimulation. The control epithelium (Fig. 6 C) received two control washes (apical and basal) and were then stimulated by forskolin after 2 h to verify that the wash protocol and extended short circuiting had no adverse effects on the cells or their forskolin response. Fig. 6 G summarizes the results from five experiments like those presented in Fig. 6 (A and B). The data demonstrate that channels blocked by benzamil at the peak of forskolin stimulation comprise the majority of channels inserted upon restimulation.

Benzamil Block Does Not Disrupt Trafficking

To evaluate the influence of benzamil on membrane trafficking, $C_T$ was recorded from cells subjected to the protocol of Fig. 6 B. Note that even when the $I_{SC}$ is almost completely blocked (Fig. 7 A), the $C_T$ response to forskolin was similar to that observed in control epithelia. This indicates that channel blockade did not influence the trafficking processes that result in an increased ENaC delivery to the cell surface in response to forskolin. This result is consistent with the lack of effect of amiloride on $C_T$ presented in Fig. 1.

Newly Synthesized ENaC Is Not Essential for the cAMP Response

To determine whether newly synthesized channels were required for repeated rounds stimulation, epithelia were treated with 200 μg/ml CHX for 30 min before the first forskolin stimulation to block protein synthesis and channel progression through the biosynthetic pathway. The efficacy of CHX disruption of protein production was determined by [$^{35}$S]Met/Cys incorporation in control and CHX-treated epithelia as described in MATERIALS AND METHODS. Phosphoimager radioactive counts for control epithelia were 13,570 ± 1,770 ($n =$
Acute ENaC Regulation by Recycling from an Intracellular Pool

6) compared with CHX-treated cells, which had only 1,120 ± 98 (n = 6), representing a reduction of ~92% in radiolabeled protein. This demonstrated that protein production was essentially eliminated by treatment with 200 µg/ml CHX. CHX remained in the bathing solutions for the duration of the experiment. The CHX-treated cells were able to respond to repetitive stimulation, as observed for control epithelia. However, the
\[ \Delta I_{SC} \] response ran down more quickly than under control conditions, resulting in a loss of \( \sim 15\% \) of \( \Delta I_{SC} \) per stimulation cycle (Fig. 8 A). From these data, it would appear that newly synthesized ENaC does not make up the majority of channels inserted into the apical membrane in response to repetitive forskolin stimulation; rather, synthesis augments the recycling pool by \( \sim 5\% \) per cycle, since the decline in \( \Delta I_{SC} \) was slightly faster after CHX treatment. The effect of prolonged CHX treatment on basal, unstimulated currents is presented in Fig. 8 B. From the fit, control CHX-untreated epithelia had a basal current half-life of \( \sim 20 \) h, whether they were repeatedly stimulated or not. In CHX-treated epithelia, the basal current decay was more rapid (current half-life \( \sim 4 \) h). The rundown in current under control conditions likely reflects fatigue due to the extended duration of the experiments. Nevertheless, by blocking channel synthesis, an indication of the functional half-life of channels inserted into the apical membrane in response to forskolin has been obtained, and these data were used to model apical ENaC recycling (see below).

**Brefeldin A Disrupts ENaC Recycling**

The effect of disrupting channel trafficking from the trans-Golgi network (TGN) to the plasma membrane was examined using BFA. A typical \( I_{SC} \) trace from a BFA-treated CCD epithelium (Fig. 9 A) demonstrates that the initial forskolin response was not impaired, but that subsequent stimulations were markedly reduced. Data obtained for the second \( I_{SC} \) response to forskolin in the presence of BFA are summarized in Fig. 9 B.

The effect of BFA was reversible, but the time taken to restore full function was dependent on the initial dose and incubation time (unpublished data). Full responsiveness was reestablished in cells receiving 5 \( \mu \)g/ml BFA after two stimulation cycles, or \( \sim 2 \) h. The effect of BFA on \( \Delta C_T \) is presented in Fig. 9 C. The diminished \( \Delta I_{SC} \) results from an inability of the cells to deliver ENaC-containing vesicles back to the apical surface since the forskolin-induced \( \Delta C_T \) response to a second stimulation was significantly reduced by BFA.

**Chloroquine Blocks ENaC Endocytosis**

The effect of inhibiting ENaC endocytosis from the apical surface was investigated by treating CCD epithelia with 100 \( \mu \)M chloroquine. Chloroquine has been used
to nonselectively inhibit endocytic pathways by alkalinization of late endosomal vesicles (Tietze et al., 1980). CCD epithelia were preincubated in chloroquine for 4 h to elicit a maximal effect. Addition of chloroquine under basal conditions produced a steady increase in $I_{SC}$ over 4 h ($105 \pm 45\%$, $n = 4$), presumably due to inhibition of channel endocytosis (see Fig. 10 C, below). Prolonged chloroquine incubation resulted in a reorganization of the endocytic compartment, as typical punctuate endosomal vesicles coalesced into larger structures (Fig. 10, A and B). The $I_{SC}$ increase suggests that apical channel density under basal conditions is determined by a constitutive turnover of channels by endocytic and exocytic trafficking pathways.

During stimulation (Fig. 10 C), forskolin was able to elicit an initial response, but the increase in $I_{SC}$ could not be reversed even with an extended wash protocol using double the wash volume. In addition, further $\Delta I_{SC}$ responses were attenuated due to the rising, poorly reversible current. These findings are summarized in Fig. 11 A, where the percent increase in $I_{SC}$ for two successive forskolin stimulations is plotted. Therefore, the lack of response to a second forskolin stimulation probably reflects impaired channel retrieval from the apical surface. In addition, chloroquine prevented the reorganization of channels into a recycling pool following stimulus removal, since the capacitance response to a second stimulus was significantly smaller than control (Fig. 11 B). This, taken together with the fluorescence images (Fig. 10, A and B), indicates that chloroquine’s effect may involve more than a collapse of pH gradients across endocytic vesicles; rather, it results in compartment rearrangement so that channels cannot be retrieved into the apical recycling pool.

Actin Is Required for ENaC Insertion, but not Retrieval

The trafficking of intracellular vesicles is closely associated with cytoskeletal elements (see reviews see Valentinj et al., 1999; Park et al., 2000; Eitzen, 2003). The effect of actin disruption on forskolin stimulation of ENaC currents was examined by selectively depolymerizing actin in CCD epithelia using LatA. The micro-
Butterworth et al.

Graphs of Fig. 12 (A–C) illustrate actin disruption induced by a 30-min treatment with two concentrations of LatA, which induced depolymerization of actin stress fibers (at 200 nM) and a breakdown of cortical actin (at 1 μM). At 1 μM LatA, most intracellular actin was depolymerized, as reflected by the reduced phalloidin staining (Fig. 12 C). This breakdown of actin stress fibers was detected electrically as a reduction in transepithelial resistance or an increase in recorded conductance from 1.2 ± 0.2 mS before LatA treatment to 7.0 ± 0.4 mS (n = 5) at the end of the experiments. The point at which this occurred is evident from the current pulse magnitudes shown in Fig. 13 A. Nevertheless, it was possible to maintain the cells under voltage clamp for 30–60 min during the LatA-induced decline in Rf (from ~2,000 to ~200 Ω/cm²), but a prolonged restimulation protocol could not be performed. When LatA was added at the peak of the forskolin response (Fig. 13 A), it was possible to washout the agonist and return to the basal prestimulated ISc; however, forskolin restimulation produced no significant ISc response. Likewise, if LatA was added before forskolin stimu-

---

**Figure 10.** Chloroquine treatment. (A and B) Fluorescence micrographs from mCCD cells transiently expressing GFP-Endo to label endocytic compartments. Typical punctuate fluorescent staining pattern of endosomes (green) in untreated cells (A) collapsed to distended coalesced structures (B) in chloroquine-treated cells (white bar = 5 μm, nuclei labeled blue). (C) Representative ISc trace from chloroquine-treated epithelium with repeated forskolin stimulation (gray bar). Note large dips in trace are the result of extended wash protocol to ensure total removal of forskolin from basal chamber. Addition of amiloride (black bar) following the third stimulation demonstrated that the majority of recorded ISc was due to ENaC.

**Figure 11.** Chloroquine treatment summary. (A) Percentage ISc increase for two forskolin stimulations (1 and 2). Control epithelia (white bar) produced ~80% increase in ISc for each stimulation with the previously noted decline in response with successive stimuli. The first ISc response for chloroquine-treated epithelia (black bar) was not statistically significantly different from control; however, ΔISC for the second stimulation was significantly reduced. (B) Percent C T increase from prestimulated value for second forskolin stimulation (number 2 in A) indicates that chloroquine-treated cells (black bar) had a significantly reduced response when compared with controls (n > 3; *, P < 0.05).
Acute ENaC Regulation by Recycling from an Intracellular Pool

These findings suggest that actin is required to shuttle ENaC to the apical surface, but that it may not be critical for subsequent ENaC endocytosis. Fig. 13C summarizes results from five experiments similar to the one presented in Fig. 13A. The first forskolin ISc response before LatA treatment is not significantly different from control cells. Following actin depolymerization, LatA-treated epithelia had a significantly reduced forskolin response when compared with control cells.

DISCUSSION

The mechanisms involved in the acute regulation of ENaC activity have been under investigation for some time in a number of model systems, such as toad urinary bladder (Coleman and Wade, 1994; Kleyman et al., 1994b; Weng and Wade, 1994), frog skin (Els et al., 1991; Lyall et al., 1994; Chou and Els, 1997; Nielsen, 1997), cultured A6 or M1 kidney cell lines (Marunaka and Eaton, 1991; Chalfant et al., 1993; Kleyman et al., 1994a; Verrey, 1994; Verrey et al., 1995; Edinger et al., 1999), and overexpression systems, and have employed a variety of physiological and biochemical methods (Reif et al., 1986; Garty and Palmer, 1997; Morris et al., 1998; Erlij et al., 1999; Weisz et al., 2000; Morris and Schafer, 2002; Rossier, 2002). The cloning of ENaC (Canessa et al., 1994; McDonald et al., 1994, 1995) offered a new array of techniques, including antibody-based labeling; however, the mechanisms responsible for the acute regulation of ENaC activity at the apical surface have remained unclear.

Two hypotheses have been proposed to account for the acute increase in Na\(^+\) transport that follows an intracellular cAMP rise, either a change in the open probability (Po) of apical membrane-resident ENaC (Oh et al., 1993; Bradford et al., 1995; Senyk et al., 1995), possibly by direct channel phosphorylation (Shimkets et al., 1998), or the recruitment of ENaC from subapical storage pools to increase channel number (N) in the apical membrane (see Rossier, 2002; Snyder, 2002). These modes of regulation need not be mutually exclusive. Nevertheless, a body of evidence has accumulated to suggest that channels are recruited from intracellular stores and are inserted into the apical membrane on cAMP stimulation (Kleyman et al., 1994a; Snyder, 2000; Morris and Schafer, 2002). The subsequent fate of these channels (i.e., degradation versus recycling), as well as the source of channels entering the apically recruited pool (i.e., recycling versus synthesis), remains uncertain. By making use of repet-

![Figure 12](image_url)
Butterworth et al.

tive cAMP stimulation of polarized mouse CCD epithelia, we have begun to address some of these issues. The data are consistent with the concept that much of the channel population that contributes to the cAMP-stimulated \( I_{sc} \) is inserted into the apical membrane from a recycling pool. In addition, a significant proportion of those channels retrieved from the apical surface following agonist washout are returned to this pool for further rounds of stimulation.

cAMP Stimulation Increases ENaC Number

The action of cAMP to increase apical ENaC number has been demonstrated previously using electrophysiological, biochemical, and imaging methods (Snyder, 2000; Butterworth et al., 2001; Morris and Schafer, 2002). These findings are further reinforced by the surface biotinylation data presented in Fig. 2. Densiometric quantitation of the Western blots demonstrated an increase in the three ENaC subunits exposed at the apical surface following forskolin stimulation. This apparent increase in channel number contributed an 83 ± 6% increase in \( I_{sc} \) and was paralleled by a 15.3 ± 1.5% increase in \( C_T \). Forskolin stimulation was reversible, as \( I_{sc} \) and \( C_T \) declined to baseline levels on removal of the cAMP agonist; however, the potential for endocytosed ENaC to be organized into a recycling pool has not been investigated from a functional perspective. The targeting of ENaC for endocytic retrieval from the apical surface has been shown to involve the binding of Nedd4-2 WW domains with the PY motifs on ENaC COOH termini. This results in channel ubiquitination, endocytosis, and possible lysosome- and proteasome-mediated degradation (Goulet et al., 1998; Harvey et al., 1999; Abriel et al., 1999; Farr et al., 2000; Staub et al., 2000; Snyder et al., 2001, 2004; Debonneville and Staub, 2004). Nevertheless, we do not know how the extent of ENaC ubiquitination correlates with its endocytic retrieval and its eventual fate (i.e., degradation versus recycling).

Intracellular Source of ENaC for Apical Insertion

By repetitively stimulating CCD epithelia up to six times, it was possible to examine the influence of reagents that disrupt channel production or vesicle traf-
ficking on the cAMP-dependent ΔI_{SC}. Repeated stimulation produced a relatively small decline (~10%) in the forskolin response with each agonist addition, as shown in Fig. 4 B. Newly synthesized channels were not required to elicit these repetitive responses, since CHX preincubation produced a qualitatively similar response profile. ENaC production did, however, contribute to the recycling pool, as ΔI_{SC} declined at a faster rate than in control cells (~15%). In a study examining repeated vasopressin stimulation of toad urinary bladder, CHX treatment did not significantly alter the first response, but an ~20–30% decline in subsequent responses was noted (Weng and Wade, 1994).

Estimation of ENaC half-life using biochemical methods is one means of obtaining information about channel lifetimes under both basal and stimulated conditions. Nevertheless, previous studies have produced conflicting results, with half-lives ranging from minutes to several hours, depending on the methods and the system studied (Valentijn et al., 1998; Weisz et al., 2000; de la Rosa et al., 2002). In addition, differences have been reported in the half-lives of individual ENaC subunits. In a study investigating endogenously expressed channels in the A6 epithelia, de la Rosa et al. (2002) calculated a half-life of ~10–18 min for channels at the apical surface using a pulse-chase protocol, whereas the intracellular channel pool exhibited a half-life of 40–80 min. Differences in half-life may be due in part to the use of different cell lines or expression systems and experimental protocols performed at 22–28°C versus mammalian systems at 37°C. However, Weisz et al. (2000) made use of the same A6 cell line and demonstrated a long-lived (>24 h) half-life for the surface-labeled channel pool, whereas the half-life of the intracellular pool was shorter with the δ-ENaC subunit half-life of only 5 h. These findings were in agreement with the observations of Kleymen et al. (2001), who found a long half-life for the αENaC subunits in A6 cells. Again in the same cell line, 35S-methionine labeling of ENaC demonstrated a short (40–50 min) half-life for each ENaC subunit (May et al., 1997). In contrast, ENaC exogenously expressed in MDCK epithelia exhibited a half-life of ~60 min for both intracellular and surface ENaC (Hanwell et al., 2002). In oocytes, a half-life of ~4 h was obtained from pulse-chase experiments (Valentijn et al., 1998). A complication in these biochemical studies potentially arises from the use of various antibodies that may not recognize the recently identified furin-cleaved forms of channel subunits attributed to intracellular ENaC processing (Hughes et al., 2003, 2004a,b).

These findings could be consistent with either rapid channel degradation (apical t_{1/2} ~15 min) or with channel recycling (apical t_{1/2} of several hours). A brief (min) apical ENaC half-life suggests that biosynthesis must be the predominant source of apically inserted channels, as most ENaC labeled at the surface would be degraded.

Nevertheless, these studies do not address the turnover of functional channels at the cell surface or their fate after endocytic retrieval. By examining ENaC currents in a polarized mammalian CCD cell line that endogenously expresses ENaC, we have collected data concerning the apical turnover of functional channels. Since currents remained fairly stable in the presence of CHX, including the ability to repeatedly stimulate I_{SC} with forskolin, our data suggests that active ENaC is fairly stable and longer lived than some previous biochemical studies suggest.

**Contribution of Recycled Channels to Apical ENaC Insertion**

The fairly persistent response to repeated stimulation could reflect a substantial intracellular storage pool of ENaC, rather than a channel recycling process. To further test the recycling hypothesis, NiCl₂ and benzamil was employed to selectively block ENaC at different times during repetitive stimulation. For NiCl₂ inhibition of the stimulated I_{SC}, a significant reduction in subsequent responses to forskolin suggested that channels irreversibly blocked by NiCl₂ were being returned to the membrane on restimulation. When channels were blocked by benzamil in the absence of agonist, the magnitude of the subsequent response to forskolin stimulation was not significantly different from that observed in untreated cells. This finding suggests that channels were inserted from a subapical pool that was not affected by benzamil block of channels at the epithelial surface.

When benzamil was added at the peak of the forskolin response, however, it was possible to elicit only small responses to restimulation (ΔI_{SC} = 12.5 ± 2.6% of control). This suggests that the majority of the channels that normally contribute to the restimulated I_{SC} response are retrieved from the apical surface after the prior stimulation. The small response to restimulation could result from dissociation of the slowly reversible blocker, or from channels delivered to the apical surface from another source, most likely the biosynthetic pathway. The results from CHX-treated cells suggest a ~5–10% contribution of channel biosynthesis to the observed ΔI_{SC} (Fig. 8), supporting this concept. Furthermore, benzamil did not affect the ΔC_{i} response to forskolin, suggesting that the trafficking mechanism that returns channels back to the apical membrane are independent of channel block.

**Properties of the Recycling Pathway, Effects of BFA, Chloroquine, and LatA**

We examined the actions on ENaC recycling of reagents that alter membrane trafficking and cytoskeletal organization using the restimulation protocol. Treatment with
BFA did not affect the first response to forskolin, but it produced a marked inhibition of the subsequent $\Delta I_{SC}$. The finding that it was possible to elicit one cycle of forskolin stimulation–recovery suggests that ENaC-containing vesicles available for channel insertion are distal to the Golgi and TGN. The response to restimulation was significantly reduced, however, indicating that ENaC is recycled through BFA-sensitive compartments. Results from a study using BFA-treated toad bladder produced similar findings in response to repeated ADH stimulation; an initial stimulation could be elicited, but subsequent stimulations were abolished by pretreatment with 5 µg/ml BFA (Weng and Wade, 1994). The effect of BFA on ENaC recycling was unexpected in view of its well-documented inhibition of AP-1–dependent trafficking in early compartments of the protein secretory pathway. Endocytic retrieval of ENaC is anticipated to rely on AP-2–mediated processes, which are BFA insensitive, but this has not been demonstrated for ENaC. However, BFA has been observed to produce a tubulation of endosomal compartments (Lippincott-Schwartz et al., 1991; Prydz et al., 1992). It is also possible that reorganization of ENaC in the subapical pool involves trafficking through ARF-1/BFA-sensitive pathways.

The effect of disrupting the endocytic pathway with chloroquine was also examined. Chloroquine is reported to de-acidify endocytic compartments, preventing traffic of plasma membrane–derived vesicles to lysosomal and other degradation pathways (Tietze et al., 1980; Jones et al., 2004). If this were the only effect of chloroquine, a relatively slow accumulation of ENaC at the apical surface from biosynthetic pathways might be expected. A recycling channel population, when mobilized, would contribute to the accumulation of apical ENaCs. In agreement with these assumptions, the baseline ENaC current increased over time, with a maximum level reached at $\sim$3–4 h (105 ± 45% increase over the initial current). It was possible to stimulate cells even at this elevated basal $I_{SC}$, but the stimulated currents could not be reversed, even by extensive washing. Studies of the cycling of Golgi proteins to the cell surface showed a chloroquine-induced redistribution of early endosomal markers into enlarged vesicles lacking compartment identity (Puri et al., 2002). Although these cells were not labeled with early endocytic compartment reporters, it is reasonable to assume from previous work that they would be redistributed into these enlarged structures (Linstedt et al., 1997; Bachert et al., 2001). Similarly, it is likely that essential endocytic trafficking proteins accumulate within these structures and are therefore not available for channel retrieval and redistribution. This disruption of the recycling compartment, together with the accumulation of ENaC at the cell surface, disturbed the normal membrane trafficking response to restimulation, which was reflected by a reduction in the forskolin-induced $C_T$ response.

The extent of actin cytoskeletal involvement in ENaC recycling was assessed by selectively depolymerizing F-actin in mCCD epithelia using LatA. Addition of LatA before forskolin diminished the $I_{SC}$ stimulation observed in control cells. A significantly reduced $I_{SC}$ response to forskolin addition was observed for all investigations of LatA-treated cells (all recordings were longer than 15 min after forskolin addition). In contrast, LatA treatment did not affect recovery from forskolin stimulation washout. These results indicate that an intact actin cytoskeleton is required for ENaC insertion, but is not required for subsequent channel endocytosis. This finding agrees with previous studies in frog skin that examined the effect of actin depolymerization on the response to vasopressin (Els and Chou, 1993). Other studies have implicated a direct actin/ENaC interaction in the modulation of channel gating and conductance, with different forms of actin added to ENaC reconstituted in lipid bilayers or coexpressed in oocytes (Jovov et al., 1999; Berdiev et al., 2001; Copeland et al., 2001). The addition of LatA either before or at the peak of the forskolin response did not alter steady-state ENaC currents, even when it was clear from the drop in transepithelial resistance that LatA had elicited actin depolymerization. We cannot, however, preclude the association of actin with ENaC in the membrane, even though apical actin staining was largely abolished.

**ENaC Recycling Model**

The concept that ENaC is acutely inserted into the apical membrane of Na$^+$-absorbing epithelia from a vesicle-based recycling pool is supported by several results emerging from the repetitive stimulation protocol described here. A model to describe ENaC insertion–recycling pathways was developed using software (Model Maker V4; Model Kinetix) to fit the time courses of $I_{SC}$ responses in control experiments, and these compartments and kinetic parameters were then tested using results from experiments where ENaC trafficking was perturbed. A schematic of the rates and corresponding half-lives included in the model under control conditions is presented in Fig. 14.

The apical membrane was described as a compartment where channels were inserted or retrieved according to rates obtained from the control data. Stimulation or retrieval events were set to occur at 30-min intervals, to match the timing of forskolin stimulation and washout periods; no attempt was made to model the transient in $I_{SC}$ observed when the basolateral chambers were washed. A value of 100 (channels) was arbitrarily assigned to the apical compartment; also, no alteration in $P_o$ or unitary current during these responses was assumed, so that the derived $I_{SC}$ trace re-
Acute ENaC Regulation by Recycling from an Intracellular Pool

The repetitive stimulation protocol under control conditions was modeled with the assumption that ENaC is derived entirely from the intracellular recycling pool. Since the average maximum \( I_{sc} \) response was 83 ± 6%, the size of the recycling pool was set to an initial value of 85. ENaC insertion was modeled as a single exponential rise to a maximum value, using insertion rate constants calculated from the control data. Recovery from stimulation was modeled as an exponential decay, again using rates calculated from the control data. In the model, all channels retrieved during a 30-min washout were placed back into the recycling compartment (less those lost to degradation or not endocytosed), to be available for insertion on the next stimulation. The rate of current decay measured in the presence of CHX in unstimulated epithelia was used to define the biosynthetic production rate of functional channels, and this, together with a degradation component, was included in the model to account for channel turnover. From these data, it was not possible to detect the presence of multiple series compartments, e.g., within the recycling pathway; therefore, the model was constructed with only two compartments, the apical membrane and the recycling pool.

Once a single control trace was obtained, channel insertion and/or retrieval rates were altered to simulate the experimental results (Fig. 15). The simulated trace illustrating changes in functional channel number with repeated stimulation over time is presented in Fig. 15 A (trace 1). The model approximates real data, but with some observable differences. First, the rate of channel insertion with each stimulation declined in the actual recordings, but this was not recapitulated in the model, since subsequent responses were initially modeled with identical parameters to that of the first. Second, the decline in \( \Delta I_{sc} \) with time can be attributed to a failure of channels to reenter the recycling pool during recovery, either due to insufficient time between stimuli or to degradative loss. However, the loss in \( \Delta I_{sc} \) in actual traces was more rapid than predicted by the model. When a loss of 2–5% of the recycling channels was included in each round of stimulation, the model approximated the actual data with reasonable accuracy (Fig. 15 A, trace 1 vs. 2). Having achieved a reasonable simulation of the control \( I_{sc} \) traces, model parameters were altered to predict changes in cell surface ENaC number following perturbations in channel production or trafficking.

To simulate CHX block of ENaC synthesis, the channel production rate was set to zero. The resulting traces approximated the data closely (Fig. 15 B), with the drop in \( I_{sc} \) due primarily to a decline in apical insertion rate. After the fourth stimulus, there was no additional response to stimulation, as the supply of channels entering the recycling pool was exhausted. Thus, although the contribution of ENaC synthesis to individual \( I_{sc} \) responses is minor relative to the recycling pathway, it becomes highly significant when there is multiple repetitive turnover of the apical pool. It was not possible to determine whether newly synthesized channels are destined primarily for the apical membrane or the recycling compartment, and additional experiments to target these pathways will be required.
To model the action of BFA, which is reported to inhibit anterograde trafficking from the TGN, the recycling channel pool was kept intact, consistent with the initial stimulation–recovery cycle that was unperturbed in BFA-treated cells. Thereafter, ENaC trafficking to the surface membrane was prevented by setting subsequent insertion rates (from both recycling and synthesis) to zero. As expected, the model produced only one response (Fig. 15 C), followed by decay of the basal current. The model accounted for the current decay due to block of synthetic delivery (as for CHX), but the presence of small subsequent responses in the actual experimental data indicates that BFA did not completely block channel insertion. Either the action of BFA was incomplete, or a model featuring insertion of only a fraction of channels in the recycling pool would provide a better description of the data. In the latter case, multiple series compartments within the recycling pathway may provide a more realistic scheme.

The concept of a channel recycling pool is supported by this compartmental model. Other hypotheses were tested but failed to generate simulations that approximated the data as well as the model of Fig. 14, but one in particular deserves comment. The concept of a large subapical pool of channels that does not reload from the apical compartment could contribute to repetitive responses; in this model, retrieved channels were targeted directly to the degradation pathway. The ISC trace predicted for this model (Fig. 15 D) shows that the forskolin response remains fairly consistent until the intracellular channel pool (set at 500) is exhausted, at which time the current responses are eliminated. The abrupt drop in $\Delta I_{SC}$ predicted by this model was not observed. Although the timing of current collapse would be determined by the initial pool size, the consistency of individual responses (lack of $\Delta I_{SC}$ decay) does not recapitulate observed traces (Fig. 15 A). Thus, although there appear to be large pools of individual channel subunits intracellularly in some systems, especially in studies where ENaC is overexpressed (Valentijn et al., 1998), a model featuring recruitment from such a pool without channel recycling does not provide the best description of these data.

Conclusions

By examining the $I_{SC}$ and $C_T$ responses to repetitive cAMP stimulation of mouse CCD epithelia, a subapical channel ENaC recycling pool has been identified. ENaC is recruited to the apical membrane from this pool on acute stimulation, and following stimulus removal, a large fraction of the endocytosed channels is returned to this recycling compartment, to be available for additional agonist responses. A simple compartmental model provides theoretical support for the presence of this ENaC recycling pool. Definition of the
number and nature of the intracellular compartments comprising the ENaC recycling pathway will require additional studies.

APPENDIX

Nyquist Plots

The epithelium was modeled in the usual manner (Fig. A1), as an equivalent circuit consisting of serial RC elements corresponding to the apical and basolateral membranes, together with the parallel, paracellular resistance. The epithelial impedance is displayed as a Nyquist plot, permitting calculation of the total epithelial capacitance. The latter is determined by the area of the individual membranes (typically 1 μF/cm²) and its dielectric properties. The transepithelial impedance ($Z_T$) of membranes in such a model has been described previously (Paunescu and Helman, 2001a) as

$$Z_T = \frac{(Z_a + Z_b)R_p}{Z_a + Z_b + R_p},$$

where $R_p$ is the paracellular (shunt) resistance (see model circuit) and $Z_a$ and $Z_b$ are the impedances of the apical and basolateral membranes, respectively. By examining current response to an imposed sinusoidal voltage, the impedance of a single membrane ($Z_m$) at a given frequency can be described as (Singh et al., 2002)

$$Z_m(f) = \frac{R_m}{1 + (R_mC_m\omega)^\alpha},$$

where $j = \sqrt{-1}$, $\alpha = 2\phi/\pi$, and $\phi$ = phase shift angle in degrees.

It was demonstrated previously that the measured capacitance is frequency dependent (Paunescu and Helman, 2001a), especially at lower frequencies, making the time constants for the RC element frequency dependent. This was verified using the present experimental system (Van Driessche et al., 1999), which records epithelial capacitance values at five frequencies simultaneously. However, all values for capacitance were obtained >4 kHz where $C_T$ values become independent of the membrane resistance, as noted above. According to the model below, $C_T$ is determined by the capacitances of the apical and basal membrane in series:

$$\frac{1}{C_T} = \frac{1}{C_a} + \frac{1}{C_b}.$$

As the calculation of $Z_m$ yields a complex number, the results are presented as a Nyquist plot, in which the magnitude and phase of the frequency response are plotted on orthogonal axes, real and imaginary. Points are plotted for each frequency in a logarithmic range of 78 frequencies in this application.

Compartmental Modeling

The modeled traces were obtained by making use of object modeling software (Model Maker V4; Model Kinetix), which readily allowed schematic compartmental models to be constructed. The model was integrated over desired time courses to produce a model trace. The recycling pool (described in discussion) and apical membrane were constructed as compartments as illustrated in Fig. A2. Rates of insertion and retrieval for acute and constitutive ENaC trafficking were calculated from recorded amiloride-sensitive ISC responses to forskolin and forskolin washout and CHX treatments, respectively. Insertion and retrieval events were modeled as single exponential rise and decays for each compartment; for example, for the first round of stimulation, channels inserted into the apical membrane from the recycling pool could be described by

$$M_t = M_0 + R \cdot (1 - e^{-k_i}), \quad (A1)$$

where $M_0 = 100$ (starting value), $R = 85$, and $k_i = 0.15$ min⁻¹.

For the first stimulus washout event, removal of channels from the apical surface to the recycling compartment would be described by
Thus, the rate of change of the apical membrane channel pool during insertion events, set to occur over 30 min, was described as insertion from the recycling pool and production, with constitutive degradation by the following differential equation:

$$\frac{dM_t}{dt} = -k_d(M) + k_i(R) + k_p(P),$$  \hspace{1cm} (A3)

with $M = 100$, $R = 85$, $P = 10,000$ (initial values), $k_i = 0.15 \text{ min}^{-1}$, $k_p = 0.0022 \text{ min}^{-1}$, $k_d = 0.0028 \text{ min}^{-1}$.

Rates of the stimulus-washout events were set to start after a 30-min insertion event and were described as removal from the apical membrane (retrieval) and loss to degradation with constitutive insertion from production as

$$\frac{dM_t}{dt} = +k_p(P) - (k_i + k_d)(M),$$  \hspace{1cm} (A4)

with $M = M_0 + \text{number of inserted channels}$ (e.g., after 30 min $M_{30} = 183$) and $k_c = 0.14 \text{ min}^{-1}$.

The authors would like to thank C. Rice and C. King for technical assistance and Drs. Kathryn Peters and Mark Knepper for donation of ENaC antibodies.

This work is supported by the National Institutes of Health grants RO1 DK54814 (to R.A. Frizzell) and RO1 DK057718 (to J.P. Johnson) and by a postdoctoral fellowship from the Cystic Fibrosis Foundation and training grant DK061296-03 (M.B. Butterworth).

Lawrence G. Palmer served as editor.

Submitted: 10 June 2004
Accepted: 7 December 2004

REFERENCES


Debonneville, C., and O. Staub. 2004. Participation of the ubiquitin-conjugating enzyme UBE2E3 in Nedd4-2-dependent regu-


