Red Fluorescent Protein pH Biosensor to Detect Concentrative Nucleoside Transport‡5

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Human concentrative nucleoside transporter, hCNT3, mediates Na+/nucleoside and H+/nucleoside co-transport. We describe a new approach to monitor H+/uridine co-transport in cultured mammalian cells, using a pH-sensitive monomeric red fluorescent protein variant, mNectarine, whose development and characterization are also reported here. A chimeric protein, mNectarine fused to the N terminus of hCNT3 (mNect.hCNT3), enabled measurement of pH at the intracellular surface of hCNT3. mNectarine fluorescence was monitored in HEK293 cells expressing mNect.hCNT3 or mNect.hCNT3-F563C, an inactive hCNT3 mutant. Free cytosolic mNect, mNect.hCNT3, and the traditional pH-sensitive dye, BCECF, reported cytosolic pH similarly in pH-clamped HEK293 cells. Cells were incubated at the permissive pH for H+-coupled nucleoside transport, pH 5.5, under both Na+-free and Na+-containing conditions. In mNect.hCNT3-expressing cells (but not under negative control conditions) the rate of acidification increased in media containing 0.5 mM uridine, providing the first direct evidence for H+-coupled uridine transport. At pH 5.5, there was no significant difference in uridine transport rates (coupled H+ flux) in the presence or absence of Na+ (1.09 ± 0.11 or 1.18 ± 0.32 mm min⁻¹, respectively). This suggests that in acidic Na+-containing conditions, 1 Na+ and 1 H+ are transported per uridine molecule, while in acidic Na+-free conditions, 1 H+ alone is transported/uridine. In acid environments, including renal proximal tubule, H+/nucleoside co-transport may drive nucleoside accumulation by hCNT3. Fusion of mNect to hCNT3 provided a simple, self-referencing, and effective way to monitor nucleoside transport, suggesting an approach that may have applications in assays of transport activity of other H+-coupled transport proteins.

Nucleosides are hydrophilic molecules that require transport proteins to mediate their movement across the plasma membrane (1). Human (h)7 nucleoside transport (NT) proteins catalyze the vectorial transport of nucleosides, using either concentrative (C) or equilibrative (E) mechanisms (2). hCNTs use either a Na+ or H+ gradient to accumulate nucleosides against their concentration gradient, whereas hENTs mediate facilitated diffusion of nucleosides down their concentration gradient (3). Nucleoside transporters also transport anti-cancer and anti-viral drugs, and cellular expression of nucleoside transporters is important in cancer therapy as well as in the treatment of cardiovascular, parasitic, and viral diseases (4, 5).

Members of the SLC28 family of concentrative nucleoside transporters (CNTs) divide into two phylogenetic subfamilies: hCNT1/2 belonging to one subfamily, and hCNT3 to the other (6–8). Cation substitution and charge/flux ratio studies suggest that hCNT1/2 couple the inward movement of nucleoside to the Na+ electrochemical gradient with a 1:1 stoichiometry, whereas hCNT3 can couple nucleoside transport to either the Na+ gradient (2 Na+:1 nucleoside) or a H+ gradient (1 H+:1 nucleoside) in the absence of Na+ (9, 10). The 2:1 coupling ratio of hCNT3 allows it to develop a trans-membrane nucleoside concentration gradient up to 10-fold higher than that of hCNT1 or hCNT2 (9, 11). At pH 5.5, hCNT3 also transports uridine in the presence of Na+ with a 2 cation:1 nucleoside stoichiometry, which raises the possibility that 1 H+ and 1 Na+ may be transported per nucleoside molecule in these conditions (9–12). Up to this point, however, there has been no direct demonstration that hCNT3 can transport H+.

Concentrative nucleoside transport has previously been investigated using the Xenopus laevis oocyte expression system and both electrophysiology (two-microelectrode voltage clamp technique) and radioisotope flux measurements (6–9, 12). Electrophysiological experiments are advantageous in that they...
measure the current induced by addition of substrate in real-time, but they are time-consuming and require specialized equipment and skills. Radioisotope flux assays measure the accumulation of radiolabeled substrate. The need for radiolabeled substrate restricts the range of permeants able to be studied. In addition, radioisotope flux assays are not done in real-time and are labor-intensive, requiring large numbers of oocytes.

An attractive alternative approach for the study of hCNT3 would be to measure pH in the immediate vicinity of its intracellular face during H⁺/nucleoside co-transport. These measurements could take advantage of the remarkable progress achieved in the development of genetically encoded fluorophores (13). Indeed, all members of the extended family of homologues and variants of the Aequorea victoria green fluorescent protein (avGFP) exhibit pH-dependent changes in their fluorescent intensity. The spectral changes that occur upon a change in pH can be intensityometric (14), excitation ratiometric (14), emission ratiometric (15), or both excitation and emission ratiometric (16). The apparent pKₐ (pKₐ', equal to the pH at which the fluorescence is half-maximal in intensity) for a specific fluorescent protein (FP) is acutely dependent on specific amino acid substitutions in close proximity to the chromophore and can range from less than 3 (17, 18) to greater than 8 (19). Variants with pKₐ' values that are relatively close to intracellular pH values (i.e. ~7.3 for the mammalian cytosol (20)) are particularly useful as genetically encoded biosensors for dynamic measurement of proton concentrations in living cells.

A major development in the area of FP technology has been the identification (21) and subsequent optimization (22, 23) of red fluorescent protein (RFP) homologues of avGFP. The first (monomeric RFP 1 (mRFP1)) (23) and second (the mFruit series) (22) generation-optimized RFPs, derived from tetrameric Discosoma RFP (21), suffer from relatively low brightness relative to other common hues of FP. For example, of the three most red-shifted second generation mFruit variants (mTangerine, mStrawberry, and mCherry) (22), the brightest (mStrawberry) has only 44% of the intrinsic brightness (proportional to the product of extinction coefficient (ε) and quantum yield (Φ)) of the popular yellow FP (YFP) Citrine (24) and 76% of the brightness of enhanced avGFP. This limitation has been partially addressed by third generation mRFPs, specifically mApple and TagRFP-T, with fluorescent brightness values on par with, or better than, that of enhanced avGFP (25).

Generally speaking, the most red-shifted RFPs derived from Discosoma RFP are relatively pH-insensitive, with the majority of variants having pKₐ' values < 5 (22, 25). A notable exception is the recently reported mApple variant with a pKₐ' of 6.5 (25). The more blue-shifted of the mFruit variants (i.e. mOrange) also have pKₐ' values of 6.5 (22). Several variants of mRFP1 with pKₐ' values > 7.5 have been previously reported (26).

Here we report the engineering of a pH-sensitive mFruit variant through multiple rounds of directed evolution by random mutagenesis. This RFP, called mNectarine, is appropriate to measure physiological pH changes in mammalian cells, because it has a pKₐ' of 6.9. We have developed a new method to measure H⁺/nucleoside co-transport in mammalian cells, which utilizes hCNT3’s H⁺ coupling characteristics and the pH sensitivity of mNectarine. We fused mNectarine to the cytosolic N terminus of hCNT3 to generate mNect.hCNT3. Fusion of the fluorescent H⁺ sensor to hCNT3 enabled measurement of pH at the intracellular surface of hCNT3, and provided insight into the mechanism of hCNT3 H⁺/uridine co-transport.

**EXPERIMENTAL PROCEDURES**

**Materials**—Synthetic DNA oligonucleotides for cloning and library construction were purchased from Integrated DNA Technologies (Coralville, IA). PCR products and products of restriction digests were purified by agarose gel electrophoresis and extraction using the GenCatch™ gel extraction kit (Epoch Biolabs, Sugar Land, TX) or QIAquick® Gel Extraction kit (Qiagen, Mississauga, Ontario, Canada). Plasmid DNA was purified from bacterial cultures, using the GeneJET™ Plasmid Miniprep kit (Fermentas, Ontario, Canada) or the HiSpeed® Plasmid Purification kit (Qiagen). Restriction endonucleases were purchased from either Invitrogen or New England Biolabs (Ontario, Canada). PNGaseF and T4 DNA ligase were from New England Biolabs. Dye terminator cycle sequencing reactions were performed using the DYEnamic ET kit (Amersham Biosciences) and were analyzed at the University of Alberta Molecular Biology Service Unit. ECL chemiluminescent reagent was from PerkinElmer Life Sciences. Ammonium persulfate was from Bioshop Canada Inc. (Burlington, Ontario, Canada). Acrylamide was from Bio-Rad. Protease inhibitor mixture tablets were from Roche Applied Sciences. Donkey anti-rabbit IgG conjugated to horseradish peroxidase (SC-2317) was from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA). Anti-RFP rabbit polyclonal antibody (ab34771) was from abcam (Cambridge, MA). Poly-l-lysine, nigericin sodium salt, uridine, 3', 6'-bis(acetylxylo)-5(6)-[[[acetyloxy]methoxy]carbonyl]-3-oxo-spiro[isobenzofuran-1(3H),9'-[H]xanthene]-2',7'-dipropanoic acid 2',7'-bis[acetylxylo]methyl] ester (BCECF-AM) and 5-([N-ethyl-N-isopropyl]amiloride (EIPA) were from Sigma-Aldrich Canada (Oakville, Canada). Platinum® Pfu DNA polymerase, Dulbeco’s modified Eagle medium, calf serum, fetal bovine serum, and penicillin-streptomycin-glutamine were from Invitrogen. Round coverslips and N,N,N',N'-tetramethylethylenediamine were from Fisher Scientific. Excitation and emission filters for fluorescence screening and photometry were purchased from Chroma Technology (Rockingham, VT). The nucleotide sequence of mNectarine has been deposited in the GenBank™ nucleotide sequence data base under accession number FJ439505.

**Mutagenesis, Library Construction, and Screening**—Mutagenesis of cDNA was performed by overlap extension PCR (23) or error-prone PCR (24). PCR products were digested with XhoI and EcoRI and ligated into pBAD/His B vector digested with the same two enzymes, and used to transform electrocompetent Escherichia coli strain DH10B (Invitrogen), which were then plated on agar plates, containing LB-AMP (Luria-Bertani medium supplemented with 0.1 mg/ml ampicillin and 0.02% w/v L-arabinose). Plates were incubated for 14 h at 37 °C prior to library screening. Our previously described imaging system (27), was equipped with a 520–550 nm bandpass filter for excitation of red fluorescence from bacterial colonies plated on a
Construction of Fusion Proteins—Mammalian expression construct mNectarine (pDEJ6) was generated by digestion of mNectarine in pBAD/His B with Xhol and EcoRI and ligation into pcDNA3.1(−) (Invitrogen). hCNT3 in the mammalian expression vector pcDNA3.1(+) was generated by digestion of hCNT3 in yeast expression vector pYPGE15 (28) with Kpn1 and EcoRI and ligation into pcDNA3.1(+) (Invitrogen). mNectarin, hCNT3 (pDEJ13) was constructed by two steps of PCR and cloning. The forward primer (5′-GGCTCAGGTCAAATGTATTAGAT-3′) included an Xhol site and the last 18 bp of hCNT3. The cDNA was generated by PCR (30 cycles of 94 °C for 30 s, 60 °C for 60 s, and 72 °C for 120 s) and inserted into the Kpn1 and Xhol sites of the mammalian expression vector pcDNA3.1(+) to generate pDEJ11. The forward primer (5′-GGCGCAAGCTTATGGTACACAGGCGG-3′) included a HindIII site and the first 19 bp of hCNT3. The reverse primer (5′-GCCATACCTTGATCGTCCATGCC-3′) deleted the stop codon of mNectarin, included a Kpn1 site, and the last 21 bp of mNectarin. The cDNA was generated by PCR (30 cycles of 94 °C for 30 s, 63 °C for 60 s, and 72 °C for 120 s) and was inserted into the HindIII and Kpn1 sites of pDEJ11 to generate pDEJ13. mNectarin.hCNT3-F563C (pDEJ20) was generated by digestion of pDEJ13 and hCNT3-F563C in oocyte expression vector pGEMHE (described previously) (29) by digestion with BamHI and EcoRV, and ligation. This introduced the F563C mutation into mNect.hCNT3. All fusion protein sequences were confirmed by sequencing (DNA Core Service Laboratory, Dept. of Biochemistry, University of Alberta).

Protein Purification and Characterization—For production of mNectarine protein, E. coli strain LMG194 was transformed with the pBAD/His B expression vector containing the FP gene of interest. A single colony was used to inoculate a 4-ml culture in LB-AMP. The culture was grown for 12 h before cells were harvested by centrifugation and lysed by French Press. Proteins were purified by nickel-nitrilotriacetic acid chromatography (Amersham Biosciences), and then dialyzed into 5 mM Tris buffer, pH 7.5. Absorption spectra were recorded on a DU-800 UV-visible spectrophotometer (Beckman Coulter, Mississauga, Ontario, Canada), and fluorescence excitation and emission spectra were recorded on a Safire2 plate reader. Fluorescence pKₐ measurements were performed by diluting the diazylated protein into a series of buffered solutions (200 mM imidazole, 200 mM citric acid, and 200 mM sodium phosphate), previously adjusted to various pHs. Quantum yields were determined using mTangerine as the reference standard (22). Protein concentrations used for calculation of extinction coefficients were determined by the BCA method (Pierce).

Photostability Measurements—For photostability measurements, microdroplets were formed by vortexing a solution of the purified protein (100 μM protein in 5 mM Tris buffer, pH 7.5) mixed with mineral oil (17, 30). A sample of this suspension (~50 μl) was sandwiched between a glass slide and a glass coverslip. The slide was imaged on an Axiointer 200M inverted fluorescence microscope (Zeiss) equipped with a 75-watt xenon-arc lamp, a 20× objective, a 510–560 nm excitation filter, a 565 nm beam splitter, a 573–648 nm emission filter, and a Retiga 2000R 12-bit cooled charged-coupled device camera (QImaging). Individual drops of protein solution considerably smaller than the field of view were identified by eye under low excitation light levels (2.5% neutral density filters). Digital image acquisition was then initiated, and the neutral density filters were removed. Collected images were processed using Image Pro (Media Cybernetics) to extract the fluorescence intensity as a function of time. Photobleaching curves were processed such that the bleaching half-time represents the time to bleach from an emission rate of 1000 photons/molecule/s to 500 photons/molecule/s (30). mTangerine was subjected to bleaching under identical conditions and used as a reference standard (22).

Tissue Culture—mNectarine, mNectarine.hCNT3, mNectarin. hCNT3-F563C, and hCNT3 constructs were expressed by transient transfection of HEK293 cells (31), using the calcium phosphate method (32). Cells were grown at 37 °C in an air/C02 (19:1) environment in Dulbeco’s modified Eagle’s medium (supplemented with 5% (v/v) fetal bovine serum, 5% (v/v) calf serum, and 1% (v/v) penicillin-streptomycin-glutamine). In experiments where fluorescence of intact HEK293 cells was monitored, HEK293 cells grown on poly-L-lysine-coated 25-mm round coverslips were transiently transfected with the appropriate cDNA.

PNGaseF Treatment and Immunodetection—HEK293 cells were transiently transfected with vector, mNectarine.hCNT3, or mNectarine.hCNT3-F563C cDNA. Cell lysates were harvested in IPB buffer (1% Nonidet P-40, 5 mM EDTA, 0.15 M NaCl, 0.5% deoxycholate, 10 mM Tris-HCl, pH 7.5). Samples (20 μg of protein) were combined with 2 μl of glycoprotein denaturing solution (0.5% SDS, 40 mM dithiothreitol) and water to make a 20-μl reaction volume. Samples were denatured by heating to 100 °C for 10 min. Reactions were made to 40 μl by addition of 4 μl of 10× G7 reaction buffer (50 mM sodium phosphate, pH 7.5), 4 μl of 10% Nonidet P-40, 1 μl of PNGaseF enzyme, and water. Samples were incubated at 37 °C for 1 h. One volume of 2× SDS-PAGE sample buffer (20% (v/v) glycerol, 2% (v/v) 2-mercaptoethanol, 4% (w/v) SDS, 1% (w/v) bromphenol blue, 150 mM Tris, 2× protease inhibitor mixture, pH 6.8) was added to each sample. Prior to analysis, samples
were heated for 5 min at 65 °C and sheared through 21- and 26-gauge needles (BD Biosciences). Samples were resolved by SDS-PAGE on 7.5% acrylamide gels (33). Proteins were transferred to polyvinylidene difluoride membranes by electrophoresis for 1 h at 100 V at 20 °C in 10% (v/v) methanol, 25 mM Tris, and 192 mM glycine (34). Polyvinylidene difluoride membranes were blocked by incubation for 1 h at 20 °C in TBST-M buffer (TBST buffer (0.1% (v/v) Tween 20, 137 mM NaCl, 20 mM Tris, pH 7.5), containing 10% (w/v) nonfat dry milk) and then incubated at 4 °C for 16–18 h in TBST-M containing 1:1,000 diluted rabbit anti-RFP polyclonal antibody. Blots were incubated for 1 h with TBST-M containing 1:5000 diluted donkey anti-rabbit IgG conjugated to horseradish peroxidase. Blots were visualized using ECL reagent and a Kodak Image Station 440CF.

Measurement of Fluorescence in Intact HEK293 Cells—HEK293 cells, grown, and transfected on 25-mm glass cover-slips were mounted in a 35-mm diameter Attofluor Cell Chamber (Molecular Probes). The chamber holds a custom-built insert, reducing the internal diameter to 13 mm and chamber volume to 0.2 ml. The chamber was placed on a Leica DMRB inverted microscope, equipped with a Photon Technologies International (PTI) D-104 microscope photometer. The light source, connected to the microscope via a fiber optic cable, was a 75-watt xenon arc lamp in a PTI DeltaScan excitation monochromator, equipped with a chopper to enable dual excitation wavelength measurements. Excitation wavelengths were set to 550 nm (when monitoring mNect), or 440 and 502.5 nm (when monitoring BCECF). Wavelengths of emitted light were selected with a cube mounted in the microscope, containing a 75-watt xenon arc lamp in a PTI DeltaScan excitation monochromator, equipped with a chopper to enable dual excitation wavelength measurements. Excitation wavelengths were set to 550 nm (when monitoring mNect), or 440 and 502.5 nm (when monitoring BCECF). Wavelengths of emitted light were selected with a cube mounted in the microscope, containing either a 570 nm beamsplitter and a 575–625 nm emission filter (mNect) or a 515 nm beamsplitter and a 522.5–547.5 nm emission filter (BCECF).

Spectral Characterization of mNectarine Expressed in HEK293 Cells—HEK293 cells, transiently transfected with mNect CDNA, were solubilized with 5% Triton X-100, 50 mM phosphate, pH 7.6, and lysates collected following centrifugation for 2 min at 18,300 × g. Lysate was added to 50 mM phosphate, pH 7.6, and excitation and emission scans were collected at respective fixed emission and excitation wavelengths 590 and 550 nm, with sample in the cuvette of a PTI fluorometer. Scans were normalized by setting peak fluorescence to 1.0.

To measure spectral characteristics of intact HEK293 cells, cells were mounted on the microscope stage, as described above, and cytosolic pH was clamped to be the same as extracellular pH by perfusion in pH clamping buffer (5 mM glucose, 140 mM KCl, 5 mM potassium gluconate, 1 mM calcium gluconate, 1 mM MgSO4, 2.5 mM sodium phosphate, 30 mM Heps), containing 20 μM nigericin, with pH set to 6.5, 7.0, and 7.5. The photometer and microscope were configured for mNect fluorescence measurements, as described above. Excitation scans from 505–565 nm were collected, and all data (mNect and vector alone transfected cells) were normalized to the peak fluorescence value found for mNect-transfected cells clamped to pH 7.5.

Calibration of Fluorescence Values for pH—To convert the fluorescence values observed with pH reporters, BCECF, and mNect, transfected HEK293 cells mounted on the microscope stage were sequentially perfused with pH clamping buffers at pH values of ~6.5, 7.0, and 7.5 (35). Fluorescence counts for BCECF or mNect were measured, and in the case of mNect fluorescence, data were corrected for photobleaching (see below). A calibration curve relating the average fluorescence at each pH to the medium pH value was fitted to a straight line by linear regression. The resulting equation for the line was used to transform fluorescence count data to intracellular pH.

**H⁺/Uridine Co-transport Activity Assay**—Transfected HEK293 cells mounted on the microscope stage, as described above, were perfused at 3.5 ml/min consecutively with Na⁺-free MBSS buffer (90 mM choline chloride (ChCl), 5.4 mM KCl, 0.4 mM MgCl2, 0.4 mM MgSO4, 5.5 mM glucose, 100 mM D-mannitol, 10 mM MES, pH 7.5), Na⁺-free MBSS buffer, pH 5.5, and Na⁺-free MBSS buffer, pH 5.5, with 0.5 μM uridine. Alternatively, cells were perfused consecutively with Na⁺-free MBSS buffer, pH 5.5, Na⁺-containing MBSS buffer (90 mM NaCl, 5.4 mM KCl, 0.4 mM MgCl2, 0.4 mM MgSO4, 5.5 mM glucose, 100 mM D-mannitol, 10 mM MES), pH 5.5, and Na⁺-containing MBSS buffer, pH 5.5, with 0.5 μM uridine. Some experiments contained 0.5 μM EIPA. At the end of each experiment, cells were subjected to pH calibration, using the nigericin/high potassium method (described above). Rates of pH change during the 20-s periods before and after addition of uridine were determined as the slope (dpH/dt) of the line fitted by the least squares method. Uridine-induced pH change was calculated as: dpH/dt after addition of uridine minus dpH/dt before uridine addition. H⁺ Flux, in units of dpH/dt, was converted to flux of proton equivalents/time by multiplying dpH/dt by the previously established intrinsic buffer capacity of HEK293 cells (36). These experiments were performed in nominally CO₂/HCO₃⁻-free conditions, so it was assumed that the CO₂ buffer capacity would be negligible.

**Photobleaching Correction**—Cells were excited as described above, and fluorescence values over time were converted to F/F₀ and fitted with an exponential decay equation of the form,

\[ Y = \text{span*exp}(-K*X) + \text{plateau} \]  

(Eq. 1)

where fluorescence starts at Span + Plateau and decays to plateau with rate constant K. The half-life is 0.69/K. The variables over five experiments were averaged, and the data for each subsequent experiment was multiplied by the average decay equation.

**Kinetics of H⁺/Uridine Co-transport**—Experiments were carried out as described under H⁺/uridine co-transport activity assay with various concentrations of uridine (0–960 μM). Transport rates were obtained as described above, and plotted in Prism 4.0. Data were fitted with an equation of the form,

\[ V = V_{\text{max}}*[S]/(K_m + [S]) \]  

(Eq. 2)

where \( V_{\text{max}} \) is the velocity (V) at maximal substrate concentration ([S]) and \( K_m \) is the substrate concentration at which the rate is half-maximal.

**Statistical Analysis**—Values are expressed ± S.E. Statistical significance was determined using an unpaired t test (Prism), with \( p < 0.05 \) considered significant.
**RESULTS**

**Engineering mNectarine**—With the goal of engineering an mRFP with improved brightness, we chose mTangerine (the Q66C/T147S/Q213L variant of mRFP1) (22) for further optimization by directed evolution. We selected mTangerine because: 1) it has the highest fluorescence quantum yield of the most red-shifted second generation mRFPs (22); 2) its absorbance profile makes it well suited as a potential fluorescence resonance energy transfer acceptor from a YFP donor; and 3) relatively little effort had been previously expended on its optimization (22). We first transferred the characteristic mTangerine mutations (M66C and Q213L) to an engineered homologue that had already been subjected to extensive directed evolution for brightness and folding efficiency (mCherry2, the K92N/K138C/K139R/S147T/N196D/T202L variant of mCherry) (supplemental Fig. S1). mCherry2-M66C/Q213L had a fluorescence hue similar to that of mTangerine, yet had intrinsic fluorescent brightness that was 1.1× that of mCherry and 1.5× that of mTangerine (Table 1). This variant was used as the template for the first of five iterative rounds of random mutagenesis and manual fluorescence-based screening of large randomized libraries (each on the order of 10⁵ variants) expressed in bacterial colonies. In each round the brightest colonies were picked, plasmids isolated, and the pool of improved FP genes used in bacterial colonies. In each round the brightest colonies were picked, plasmids isolated, and the pool of improved FP genes used as the template for the subsequent library. We have successfully applied this basic strategy to the optimization of a teal FP (27), blue FPs (17), and a YFP with violet excitation (37).

During this process, the protein accumulated four additional substitutions relative to the initial template: F91L, M141V, Y151H, and K162M. The resulting protein was an mRFP with intrinsic brightness (Table 1) that was considerably greater than that of mCherry and mTangerine (22) (1.6× and 2.3×, respectively) but still somewhat less than that of mApple and TagRFP-T (25) (0.71× and 0.78×, respectively). The excitation and emission maxima of the final variant (excitation maximum at 558 nm and emission maximum at 578 nm) were different enough from that of mTangerine, and every other variant in the existing palette, that we decided to designate it with the name “mNectarine” (supplemental Fig. S1).

**pH Sensitivity of mNectarine**—Characterization of mNectarine revealed that its fluorescence was remarkably pH-sensitive, with an apparent pKₐ of 6.9 (Fig. 1, A and B). The data points shown in Fig. 1B were fitted to a curve of the general form,

\[ I_{\text{pH}} = \frac{I_{\text{max}}}{\frac{1}{10^{(pK_a - pH)}} + 1} \]  

(Eq. 3)

where \( I_{\text{pH}} \) is the fluorescence intensity at a given pH, and \( I_{\text{max}} \) is the highest fluorescence intensity at any pH. An equation of similar form was used to fit the absorbance data points as shown in Fig. 1D.

Absorbance spectra of mNectarine recorded at various pHs revealed a complex change in profile that appears to include a mixture of anionic cyan-absorbing (\( \lambda_{\text{max}} = 489 \) nm) and orange-absorbing (\( \lambda_{\text{max}} = 558 \) nm) forms of the protein at high pH and a mixture of protonated violet-absorbing (\( \lambda_{\text{max}} = 387 \) nm) and blue-absorbing (\( \lambda_{\text{max}} = 453 \) nm) forms of the protein at low pH (Fig. 1C). A reasonable explanation of these changes is that the purified and aged protein exists as a mixture of an orange-absorbing/red-fluorescing species and a “dead-end” cyan-absorbing/non-fluorescent species with an avGFP-type chromophore. Accordingly, the cyan and violet species may represent two ionization states of one distinct form of an avGFP-type chromophore, whereas the orange and blue species represent two ionization states of a Discosoma-type (or possibly an mOrange-type, containing a third heterocycle (38)) chromophore. To investigate this hypothesis, we performed a multiple linear regression analysis of the spectra acquired at each pH to examine the relative contribution of each of these four components. Efforts to fit the data with fewer than four components resulted in poor fits of the experimental data. This analysis revealed that pH-dependent changes in the absorbance of the violet (decreasing with increasing pH) and cyan species (increasing with increasing pH) occur with pKₐ values of ~8.1 (Fig. 1D), an observation consistent with the conclusion that these are indeed different ionization states of the same molecular species. The multiple regression analysis did not provide support for correlated changes in the orange and blue species as a function of pH, with the intensity of blue species remaining relatively constant as the intensity of orange species changed considerably. Two possible explanations for this discrepancy are that either the blue peak represents a distinct dead-end species, or that our data are simply insufficient to enable separation of the overlapping contributions of the blue and orange peaks.

**mNectarine.hCNT3 Fusion Proteins**—We fused mNectarine to the N terminus of the human nucleoside transporter, hCNT3. The N terminus was chosen for the fusion, because the C terminus is extracellular (8), and our goal was to measure pH at the intracellular surface of hCNT3. We wanted to use a specific and localized probe of [H⁺] because hCNT3 has a relatively low turnover rate (a value of 34 s⁻¹ for hCNT3 was found in voltage-clamped oocytes at −150 mV (39)). Because the rest-
ing membrane potential of mammalian cells is ~40 mV, a turnover rate closer to 10 s\(^{-1}\) is to be expected.

Expression of mNect.hCNT3 fusion proteins was assessed on immunoblots (Fig. 2). hCNT3-F563C, a functionally inactive mutant (29), was fused to mNectarine to serve as a negative control (called mNect.hCNT3-F563C). HEK293 cells were transiently transfected with vector, mNect.hCNT3, or mNect.hCNT3-F563C cDNA. Lysates from the cells were treated with or without PNGaseF (an enzyme that removes N-linked glycosylation from proteins (40)). Samples were immunoblotted and probed with an anti-RFP antibody. hCNT3 alone migrated as a 75 kDa band (12), whereas mNectarine was 27 kDa. Untreated mNect.hCNT3 was detected as multiple bands around 100 kDa (Fig. 2). Upon treatment with PNGaseF, only a single band was detected for the two fusion proteins (Fig. 2). The multiple banding pattern is attributed to unglycosylated, core glycosylated, and mature glycosylated bands. This is supported by the prediction that there are four putative glycosylation sites in the C terminus of hCNT3, and both hCNT1 and hCNT3 exhibit multiple glycosylation states (12, 41, 42). The presence of different glycoforms of hCNT3 will not affect the function of the protein at the plasma membrane, because hCNT3 with all glycosylation sites absent is fully functional (12).

Spectral Characterization of mNectarine Expressed in HEK293 Cells—Fluorescence excitation and emission scans were obtained for lysates of mNect-transfected HEK293 cells (Fig. 3A). The peak fluorescence excitation wavelength was 558 nm, and the peak emission wavelength was 573 nm. Fluorescence excitation spectra were collected for intact mNect-transfected HEK293 cells, or vector-alone transfected cells clamped at pH 7.5, 7.0, and 6.5 with nigericin/high potassium (Fig. 3B). mNectarine fluorescence decreased with decreasing pH (Fig. 3B), in a manner similar to that seen with purified mNectarine (Fig. 1). In intact HEK293 cells the peak excitation wavelength was 558 nm at pH 6.5 and 7.0, and 557 nm at pH 7.5 (Fig. 3B). The observed fluorescence of mNect-transfected cells can be attributed to mNect fluorescence, because little fluorescence was observed when similar numbers of vector-alone transfected HEK293 cells were subjected to fluorescence excitation scans (Fig. 3B, lower curves). A plot of normalized fluorescence in intact mNect-transfected cells versus medium pH revealed a linear relationship over the pH 6.5–7.5 range (Fig. 3C). This indicates that mNect fluorescence, upon appropriate calibration, can be used as a reporter of cytosolic pH in intact cells.

Correction for mNectarine Photobleaching—Extended periods of illumination of mNect, under pH-clamped conditions, resulted in a continuous decrease of fluorescence, consistent with photobleaching (Fig. 4A). Examination of the kinetics of
bleaching revealed they could be fitted by an exponential decay equation, which provides the basis for a method to correct for photobleaching. Under pH-clamped conditions mNect should report a constant fluorescence value, in the absence of photobleaching. The ability to correct photobleaching, by applying an exponential correction, was applied to pH 7.0-clamped cells (Fig. 4, A and B). The data revealed that the continuous reduction in signal was substantially corrected, but not fully eliminated. To limit the effects of photobleaching and the need to apply corrections, in further experiments, steps were taken to minimize the time period during which mNect was illuminated.

The ability for mNect to report cytosolic pH accurately, following photobleaching correction, was assessed by clamping cytosolic pH to medium pH, using the nigericin/high potassium method (Fig. 4, C and D). In these experiments pH was clamped sequentially to values near 7.0, 6.5, 7.5, and 7.0 (reported in Fig. 4, C and D, in the black bar) by perfusion with media of known pH. To calibrate fluorescence to cytosolic pH, a standard curve was generated, relating average fluorescence to media pH at each of the last three solutions. The average pH value during perfusion with each solution was then calculated (reported mNectarine and hCNT3 cDNA, or mNect.hCNT3, or with hCNT3 alone. These latter cells were loaded with BCECF, by incubation with BCECF-AM. Cells were perfused with nigericin/high potassium solutions at pH 7.5, 6.5, and 7.0. As described above, the average fluorescence values at each medium pH were then used to construct a standard curve, which was then applied to the fluorescence data, to transform it to intracellular pH (Fig. 5). Comparison, between the pH of perfusion medium to reported cytosolic pH determined from fluorescence, reveals that mNect, mNect.hCNT3, and BCECF are similar in their ability to report cytosolic pH. We then measured the absolute difference in pH between the medium pH and the pH reported by BCECF and mNect.hCNT3 to provide a quantitative comparison of the reliability of the two fluorescent pH reporters. For BCECF the mean absolute difference was 0.013 ± 0.003 pH units (n = 12) and for mNect.hCNT3 the mean absolute difference was 0.024 ± 0.005 pH units (n = 24). We conclude that BCECF reports on cytosolic pH more accurately than mNect.hCNT3, but that mNect.hCNT3 can be expected to report on cytosolic pH within 0.04 pH units of the “true” cytosolic pH over the pH 6.5–7.5 range.

FIGURE 3. Spectral characterization of mNectarine expressed in HEK293 cells. HEK293 cells were transiently co-transfected with cytosolic pH reporters. For BCECF the mean absolute difference was 0.038 pH units (n = 24). Together these data reveal that after correction for photobleaching, mNect, calibrated by sequential perfusion with nigericin/high potassium media in the pH 6.5–7.5 range is able to report accurately on cytosolic pH.

Measurement of Cytosolic pH by mNect, mNect.hCNT3, and BCECF—To test the reliability of mNect and mNect.hCNT3 as reporters of cytosolic pH, we compared their ability to report cytosolic pH to data collected using the well established pH-sensitive dye, BCECF (Fig. 5). HEK293 cells were transiently co-transfected with cytosolic...
Intracellular pH upon stimulation of H\(^{+}\)/nucleoside co-transport—We next examined the ability of the mNect.hCNT3 fusion proteins to report on changes in intracellular pH upon stimulation of H\(^{+}\)/nucleoside co-transport. We used uridine as the nucleoside of choice because uridine elicits the largest nucleoside-induced inward currents under Na\(^{+}\)-free, pH 5.5 conditions, which were previously found necessary for robust H\(^{+}\)/uridine co-transport (10). HEK293 cells were transiently transfected with cDNAs encoding mNect.hCNT3 or mNect.hCNT3-F563C (Fig. 6). Cells were perfused consecutively with Na\(^{+}\)-free MBSS buffer, in which ChCl was substituted for NaCl, at pH 7.5, Na\(^{+}\)-free MBSS buffer, pH 5.5, and Na\(^{+}\)-free MBSS buffer, pH 5.5, containing 0.5 mM uridine. Each experiment was then calibrated for pH using the nigericin/high potassium method (35). Co-transport of uridine with H\(^{+}\), which was initiated by the addition of uridine, was monitored by measuring intracellular pH (pH\(_i\)). Fluorescence data were corrected for photobleaching, and corrected fluorescence data were converted to pH\(_i\). Transport rates were determined by linear regression of the initial acidification rate from the first 20 s after addition of uridine minus the rate before addition of uridine.

**FIGURE 4. Correction for mNectarine photobleaching.** HEK293 cells were transiently transfected with free cysotic mNectarine cDNA, and fluorescence was monitored (λ\(_{em} = 550\) nm and λ\(_{ex} = 573\) nm). A and B, cells were perfused with nigericin/high potassium solution, pH 7.0, to clamp intracellular pH. A, raw mNectarine fluorescence at pH 7.0, declining as a result of photobleaching. B, mNectarine fluorescence at pH 7.0. Data from panel A has been corrected for photobleaching, using the approach described in methods. C and D, HEK293 cells transfected with mNect cDNA were perfused consecutively with nigericin/high potassium solutions at pHs indicated in the black bars above the curves. To minimize the period of sample illumination, samples were perfused without illumination for 480 s in each solution prior to collecting fluorescence data. Above or below each trace are the average pH values for each perfusion interval, calculated from the fluorescence data. C, data not corrected for photobleaching. D, data from panel C corrected for photobleaching, using the approach described under “Experimental Procedures.”

H\(^{+}\)/Uridine Co-transport—We next examined the ability of the mNect.hCNT3 fusion proteins to report on changes in intracellular pH upon stimulation of H\(^{+}\)/nucleoside co-transport. We used uridine as the nucleoside of choice because uridine elicits the largest nucleoside-induced inward currents under Na\(^{+}\)-free, pH 5.5 conditions, which were previously found necessary for robust H\(^{+}\)/uridine co-transport (10). HEK293 cells were transiently transfected with cDNAs encoding mNect.hCNT3 or mNect.hCNT3-F563C (Fig. 6). Cells were perfused consecutively with Na\(^{+}\)-free MBSS buffer, in which ChCl was substituted for NaCl, at pH 7.5, Na\(^{+}\)-free MBSS buffer, pH 5.5, and Na\(^{+}\)-free MBSS buffer, pH 5.5, containing 0.5 mM uridine. Each experiment was then calibrated for pH using the nigericin/high potassium method (35). Co-transport of uridine with H\(^{+}\), which was initiated by the addition of uridine, was monitored by measuring intracellular pH (pH\(_i\)). Fluorescence data were corrected for photobleaching, and corrected fluorescence data were converted to pH\(_i\). Transport rates were determined by linear regression of the initial acidification rate from the first 20 s after addition of uridine minus the rate before addition of uridine.

pH 5.5 Na\(^{+}\)-free MBSS medium induced a slow acidification of mNect.hCNT3 and mNect.hCNT3-F563C-transfected cells (Fig. 6, A and B), which is addressed below. The rate of acidification increased significantly upon addition of 0.5 mM uridine, only in mNect.hCNT3-expressing cells (Fig. 6, A and B), consistent with hCNT3-mediated H\(^{+}\)-coupled uridine transport. Quantification of the change in the rate of acidification (proton flux in units of mmol·min\(^{-1}\)) upon switching to uridine-containing medium revealed that cells transfected with mNect.hCNT3 had a proton flux of 1.08 ± 0.10 mmol·min\(^{-1}\), whereas cells transfected with the inactive hCNT3 mutant fusion protein, mNect.hCNT3-F563C, had a proton flux of −0.02 ± 0.1 mmol·min\(^{-1}\) (Fig. 6C). Assuming a H\(^{+}\)-nucleoside coupling stoichiometry of 1:1, as found for hCNT3 uridine transport when expressed in Xenopus laevis oocytes (9, 10), these data imply that the rate of change of cytosolic concentration of uridine is the same as the rate of change of [H\(^{+}\)], found using mNect.hCNT3. Together these results demonstrate that the mNect.hCNT3 fusion protein reports on H\(^{+}\)/uridine co-transport by sensing changes in pH\(_i\), associated with coupled H\(^{+}\)/uridine influx.

One caveat, however, relates to the calculation of proton flux, using the established buffer capacity of the bulk cytosol of HEK293 cells. We measured intracellular pH using mNect.hCNT3, anchored at the surface of the plasma membrane. If the buffer capacity at the membrane surface is constant; transport rates can therefore be reliably compared between different cell samples, as the buffer capacity would be the same between them. Further, it is unlikely that the buffer capacity at the plasma membrane surface differs dramatically from bulk cytosol.

Because point mutations can sometimes impair trafficking of membrane proteins, we assessed the plasma membrane localization of mNect.hCNT3 compared with mNect.hCNT3-
mNect.hCNT3 H\(^{+}\)/Uridine Co-transport

FIGURE 5. Measurement of intracellular pH, using cytosolic mNectarine, mNect.hCNT3, and BCECF. HEK293 cells were transiently co-transfected with cytosolic mNectarine and hCNT3 cDNA (A) or mNect.hCNT3 cDNA (B) or hCNT3 cDNA (C). hCNT3-transfected cells were incubated with the pH-sensitive dye BCECF-AM (C). Cells were perfused with nigericin/high potassium calibration solutions at pH values indicated in the black bars above the curves to calibrate fluorescence to pH. To minimize the period of sample illumination, samples were perfused without illumination for 480 s in each solution prior to collecting fluorescence data. Average pH values for each perfusion interval (calculated from the fluorescence data) are indicated above or below the trace. A and B, mNect fluorescence (\(\lambda_{ex} = 550\,\text{nm} \) and \(\lambda_{em} = 573\,\text{nm}\)) was corrected for photobleaching. C, BCECF fluorescence was monitored at \(\lambda_{ex} = 440\,\text{nm}\) and \(\lambda_{em} = 528.7\,\text{nm}\). F563C. HEK293 cells transfected with either mNect.hCNT3 or mNect.hCNT3-F563C cDNA had similar degrees of plasma membrane localization on the basis of imaging mNectarine fluorescence by confocal microscopy (data not shown).

To assess the reliability of the mNect.hCNT3 fusion as an assay for H\(^{+}\)/nucleoside co-transport, we measured the \(K_m\) of mNect.hCNT3 for uridine. HEK293 cells were transiently transfected with mNect.hCNT3 cDNA, and were consecutively perfused with Na\(^{+}\)-free MBSS buffer at pH 7.5, Na\(^{+}\)-free MBSS buffer, pH 5.5, and Na\(^{+}\)-free MBSS buffer, pH 5.5, containing varying amounts of uridine, from 0 to 960 \(\mu\text{M}\). The change in the rate of acidification (proton flux in units of mM min\(^{-1}\)) was quantified upon switching to uridine-containing medium at each concentration of uridine. Fig. 7 shows a representative example of three separate experiments used to calculate the \(K_m\) for uridine transport in HEK293 cells, as measured with mNect.hCNT3. The \(K_m\) determined was 72 ± 24 \(\mu\text{M}\) (\(n = 3\)), which lies between the two previously published values of 110 ± 10 \(\mu\text{M}\) and 62.4 ± 5.4 \(\mu\text{M}\) found in oocytes (10, 12). We conclude that measurement of uridine flux by following H\(^{+}\) movement with mNect.hCNT3 is an accurate assay of H\(^{+}\)/nucleoside co-transport.

Radiolabeled substrate uptake studies in oocytes found that uridine is transported to a similar extent in both acidic Na\(^{+}\)-free and acidic Na\(^{+}\)-containing conditions (10, 12). Charge/uptake experiments suggested that in Na\(^{+}\)-containing buffer, pH 5.5, both Na\(^{+}\) and H\(^{+}\) contribute to the driving force, and that one of the two Na\(^{+}\) binding sites is shared by H\(^{+}\) (10). To determine whether a H\(^{+}\) is co-transported with uridine in acidic Na\(^{+}\)-containing medium, HEK293 cells were transiently transfected with mNect.hCNT3 or mNect.hCNT3-F563C cDNA: Cells were perfused with Na\(^{+}\)-free MBSS buffer, pH 5.5, in which NaCl was replaced by ChCl, Na\(^{+}\)-containing MBSS buffer, pH 5.5, and Na\(^{+}\)-containing MBSS buffer, pH 5.5, containing 0.5 mM uridine. The Na\(^{+}\)/H\(^{+}\) exchange inhibitor, EIPA (5 \(\mu\text{M}\)), was present in all perfusion buffers. The data were collected and analyzed as described above.

Addition of uridine induced acidification in both Na\(^{+}\)-free (Fig. 8B) and Na\(^{+}\)-containing medium (Fig. 8A) in mNect.hCNT3-transfected cells. Cells expressing catalytically inactive mNect.hCNT3-F563C had no increase in acidification rate upon addition of uridine to the medium (Fig. 8C). Quantification of the increase in acidification rate following addition of uridine (Fig. 8D) revealed that Na\(^{+}\) had no effect on the acidification rate (1.09 ± 0.11 or 1.18 ± 0.32 mM min\(^{-1}\), in the absence or presence of Na\(^{+}\), respectively). This provides support for the proposed mechanism that in acidic Na\(^{+}\)-containing conditions, 1 Na\(^{+}\) and 1 H\(^{+}\) are transported per uridine molecule, while in acidic Na\(^{+}\)-free conditions, 1 H\(^{+}\) alone is transported per uridine molecule (9–12).
In H9261 medium, hCNT3 fluorescence was monitored at 20508 JOURNAL OF BIOLOGICAL CHEMISTRY.

Error bars represent standard error (n = 5). Asterisk indicates significant difference (p < 0.0001).

FIGURE 6. hCNT3 mediated H⁺/uridine co-transport measured by mNectarine-hCNT3. HEK293 cells were transiently transfected with mNect.hCNT3 cDNA (A) or with mNect.hCNT3-F563C cDNA (B). mNect. hCNT3 fluorescence was monitored at λ_em = 550 nm and λ_exc = 573 nm. Cells were perfused consecutively with Na⁺-free MBSS buffer, pH 7.5 (open bar), in which NaCl is replaced by choline chloride, Na⁺-free MBSS buffer, pH 5.5 (gray bar), and Na⁺-free MBSS buffer with 0.5 mM uridine, pH 5.5 (black bar). At the end of each experiment photobleaching-corrected fluorescence values were calibrated to pH via the nigericin/high potassium technique (not shown in figure) (35). A, pH transients observed for mNectarine.hCNT3-transfected cells. B, pH transients observed for mNect.hCNT3-F563C-transfected cells. C, quantification of the change in rate of acidification (proton flux in units of mM/min) upon switching to uridine-containing medium. Error bars represent standard error (n = 5). Asterisk indicates significant difference (p < 0.0001).

FIGURE 7. Kinetics of H⁺/uridine co-transport in HEK293 cells. HEK293 cells were transiently transfected with mNect.hCNT3 cDNA. mNect.hCNT3 fluorescence was monitored at λ_em = 550 nm and λ_exc = 573 nm. Cells were perfused consecutively with Na⁺-free MBSS buffer, pH 7.5, in which NaCl is replaced by choline chloride, Na⁺-free MBSS buffer, pH 5.5, and Na⁺-free MBSS buffer, pH 5.5, containing varying concentrations of uridine from 0 to 960 μM. At the end of each experiment pH was calibrated via the nigericin/high potassium technique, and fluorescence values were calibrated to pH. The change in the rate of acidification (proton flux in units of mM/min⁻¹) was quantified upon switching to uridine-containing medium at each concentration of uridine. This graph is a representative example of three separate experiments used to calculate the K_m of mNect.hCNT3 for uridine.

DISCUSSION

We utilized hCNT3’s H⁺-coupling characteristics to develop a new method to assay nucleoside transport in cultured mammalian cells. This assay was made possible by engineering a new pH-sensitive red fluorescent protein, mNectarine. mNectarine, fused to the N terminus of hCNT3, was able to reliably report on changes in pH at the intracellular surface of hCNT3 in real-time. Addition of uridine to mNect.hCNT3-expressing cells, in both Na⁺-free and Na⁺-containing conditions elicited an increase in the rate of acidification that was not seen in the negative control (mNect.hCNT3-F563C-expressing cells). Our data are consistent with hCNT3-mediated Na⁺/H⁺/nucleoside co-transport under acid conditions.

Using mNect.hCNT3 to measure uridine flux by monitoring H⁺ co-transport is advantageous, because it enables direct measurement of changes in intracellular H⁺ concentration, which, up until now, had only been inferred from H⁺ activation of [%]uridine influx and pH-dependent uridine-evoked currents in oocytes (8, 9, 29). The present experiments therefore provide the first direct evidence that hCNT3 transports H⁺. Because we did not directly measure uridine flux in these experiments, it is formally possible that the mNect.hCNT3 assay measures an hCNT3-mediated H⁺ flux unrelated to uridine transport. We consider this unlikely, because under similar extracellular acid medium conditions uptake of radioactive nucleoside was observed in oocytes (9, 10).

We assessed the reliability of mNect.hCNT3 as a reporter for hCNT3 H⁺/nucleoside co-transport, by measuring the K_m of mNect.hCNT3 for uridine. The K_m determined here (72 ± 24
mNect.hCNT3 ΔpH/Uridine Co-transport

mNect.hCNT3 can be used to measure the transport rate for any nucleoside or nucleoside drug that is co-transported with H⁺, so any range of substrates can be assayed for transport, even if a radioactive analogue is unavailable. This characteristic, along with utilizing a pH-sensitive fluorescent reporter, opens up the possibility of high throughput assays in which a wide range of possible substrates could be added to mNect.hCNT3-transfected cells grown in multiwell plates while monitoring fluorescence changes over time with a multiwell plate fluorometer. mNect.hCNT3 also provides the possibility of high throughput screening for inhibitors, as no high affinity CNT inhibitors have yet been identified. In contrast to hCNT3 assays performed using X. laevis oocytes (9, 10, 29), the mNect.hCNT3 assay

μM) was between two previously published values for hCNT3 expressed in oocytes (110 ± 10 and 62 ± 5 μM) (10, 12), indicating that mNect.hCNT3 provides an accurate method to measure hCNT3 kinetics.

This study illuminates the mechanism of nucleoside transport under conditions of extracellular acid and Na⁺. Previous investigations in X. laevis oocytes demonstrated that uridine was transported with high efficiency in both Na⁺-containing medium at pH 5.5, and Na⁺-free medium at pH 5.5, although to a somewhat lesser extent under Na⁺-free conditions (10, 12). hCNT3 co-transport of nucleoside with Na⁺ has been verified outside oocytes, using yeast and mammalian cells (28, 41, 46). In the current study, there was H⁺ movement in both acidic Na⁺-containing and acidic Na⁺-free medium, with no significant difference in transport rate between the two conditions. This result indicates that H⁺ is co-transported with uridine in acidic Na⁺-containing buffer. Charge/flux ratio experiments in oocytes revealed that hCNT3 functions with 2:1 cation:uridine stoichiometry in both acidic and alkaline Na⁺-containing medium, compared with a 1:1 cation:uridine stoichiometry in acidic Na⁺-free medium (10, 12). Because we find similar rates of H⁺ movement in the presence and absence of Na⁺, the present data support a model in which 1 H⁺ and 1 Na⁺ are transported per uridine molecule in acidic Na⁺-containing medium, compared with 1 H⁺ per uridine molecule in the absence of Na⁺, and 2 Na⁺ per uridine molecule in alkaline conditions.

FIGURE 8. Effect of Na⁺ on H⁺/uridine co-transport. HEK293 cells were transiently transfected with mNectarine. hCNT3 cDNA (A and B) or with mNect.hCNT3-F663C cDNA (C). mNect.hCNT3 fluorescence was monitored at λex = 550 nm and λem = 573 nm. Cells were perfused with MBSS buffer, pH 5.5 (gray bars), or MBSS buffer, pH 5.5, containing 0.5 mM uridine (black bars). Perfusion with Na⁺-containing MBSS buffer is indicated by hatched bars, and perfusion with Na⁺-free MBSS buffer, in which NaCl is replaced by choline chloride (ChCl), is indicated by un-hatched bars. Open bar indicates perfusion with Na⁺-free MBSS, pH 7.5. All buffers contained 5 mM EIPA. D, quantification of the change in rate of acidification (proton flux in units of mM min⁻¹) upon switching to uridine-containing medium. Error bars represent standard error (n = 3). N.S. indicates non-significant difference (p = 0.81).
to discern an intermediate conformation where both \( \text{Na}^+ \) and \( \text{H}^+ \) are bound.

The concentrative nucleoside transporters are primarily expressed in epithelial cells where they are involved in absorption, secretion, distribution, and elimination of nucleosides and nucleoside analogs (2, 3, 11). hCNT3 is especially abundant in the kidney, and may be the key CNT at the apical surface of the proximal tubule (3, 11). The ability of hCNT3 to couple nucleoside movement to either a \( \text{Na}^+ \) and/or \( \text{H}^+ \) gradient may be advantageous in the kidney, because the lumen of the proximal tubule is acidic, and hCNT3 could take advantage of the \( \text{H}^+ \) gradient to maximize its transport rate in these conditions. Both hCNT3 and SGLT1, which have similar transport kinetics, are present in the intestine, which also has an acidic luminal environment, particularly in the more proximal regions (48). Thus, the \( \text{H}^+ \)-coupling characteristic of hCNT3 may be physiologically and pharmacologically important (3, 9).

The use of pH-sensitive fluorescent protein fusions as an assay of transport activity extends beyond hCNT3. mNectarine could be fused to any transporter that induces a change in intracellular \( \text{pH} \). Such transporters could include the \( \text{H}^+ \)-coupled PepT1/2 peptide transporters (49, 50), SGLT1 (48), and the MCT1 monocarboxylate transporter (51), or any of the transport proteins involved in the regulation of \( \text{pH}_i \), including \( \text{Cl}^-/\text{HCO}_3^- \) exchangers (52), NHEs (53), or sodium bicarbonate co-transporters (54). PepT1 (55), MCT1 (56), NHE1 (53), and the SLC4 family members (including \( \text{Cl}^-/\text{HCO}_3^- \) exchangers and sodium bicarbonate co-transporters) (54), all have at least one intracellular terminus, so an mNectarine fusion could be constructed. SGLT1 does not have intracellular \( \text{N} \) or \( \text{C} \) termini, but FP fusions could possibly be made in the large cytosolic loop (57). Fusion of mNectarine to these proteins would report on changes in \( \text{pH} \) local to the transporter and would be more specific than simply measuring \( \text{pH} \) changes in the bulk cytosol with a \( \text{pH} \)-sensitive fluorescent protein, mNectarine, to the N terminus of hCNT3. mNectarine is an approach that could be extended to high throughput assays and could be used to report on the transport activity of any \( \text{pH} \) transporter. Taken together, our findings demonstrate that mNect.hCNT3 is a valid reporter of \( \text{H}^+/\text{nucleoside} \)-co-transport, and support a transport mechanism where \( \text{H}^+ \) and \( \text{Na}^+ \) are co-transported with uridine in acidic \( \text{Na}^+ \)-containing conditions.

Acknowledgments—The cDNA for mCherry2 was kindly provided by Nathan C. Shaner and Roger Y. Tsien (University of California, San Diego).

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