RESEARCH PAPER

hZip1 (hSLC39A1) regulates zinc homoeostasis in gut epithelial cells

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Abstract Zinc is an essential trace element required for enzyme catalysis, gene regulation and signal transduction. Zinc absorption takes place in the small intestine; however, the mechanisms by which cells accumulate zinc are not entirely clear. Zip1 (SLC39A1) is a predicted transmembrane protein that is postulated, but not conclusively proven to mediate zinc influx in gut cells. The aim of this study was to investigate a role for hZip1 in mediating zinc uptake in human enterocytes. Both hZip1 mRNA and protein were detected in human intestinal tissue. In nondifferentiated Caco-2 human gut cells, hZip1 was partially localised to the endoplasmic reticulum. In contrast, in differentiated Caco-2 cells cultured in extracellular matrix, the hZip1 protein was located in proximity to the apical microvilli. Lack of surface antibody binding and internalisation indicated that hZip1 was not present on the plasma membrane. Functional studies to establish a role for hZip1 in cellular zinc accumulation were carried out using 65 Zn. In Caco-2 cells harbouring an $hZip1$ overexpression construct, cellular zinc accumulation was enhanced relative to the control. Conversely, Caco-2 cells with an $hZip1$ siRNA construct showed reduced zinc accumulation. In summary, we show that the Caco-2 cell differentiation endorses targeting of hZip1 to a region near the apical domain. Given the absence of hZip1 at the apical plasma membrane, we propose that hZip1 may act as an intracellular sensor to regulate zinc homoeostasis in human gut cells.

Keywords Zinc homoeostasis - Zinc transporter hZip1 (SLC39A1) - Human gut

Introduction

The gut plays a key role in Zn homoeostasis through the absorption of exogenous Zn and the excretion of both endogenous and exogenous Zn. The mechanisms by which the gut maintains tight homoeostatic control of Zn are not clear, but may involve members of two major families of putative Zn transporters, the ZnT (SLC30) and the Zip (SLC39) families. The hZip4 Zn transporter is implicated in Zn uptake in the human enterocyte as its gene is mutated in the disorder acrodermatitis enteropathica, a condition where Zn absorption in the intestine is impaired, leading to depletion of body Zn stores and characteristic acral and circumorificial dermatitis, diarrhoea and alopecia (Kury et al. [2002](#page-10-0); Ackland and Michalczyk [2006](#page-10-0)). Other putative Zn transporters detected in the human gut include hZip1, hZip4 and members of the SLC30A family including ZnT1 and ZnT5 (Kury et al. [2002](#page-10-0); Cragg et al. [2005](#page-10-0)). Similar Zn transporters are present in the rodent gut (Jou et al. [2009](#page-10-0)).

The cellular localisation of Zip1 varies between different cells. In human adherent cells, a cytoplasmic or vesicular localisation of Zip1 (SLC39A1) has been described for PC3, HepG2 and COS-7, with partial overlap with BIP, a marker for the ER in COS-7 (Milon et al. [2001](#page-11-0)). A cytoplasmic location of Zip1 was reported in osteoclasts (Khadeer et al. [2005](#page-10-0)) and in human mesenchymal stem cells (Tang et al. [2006](#page-11-0)), while tagged hZip1 showed strong labelling in the vicinity of the plasma membrane of non-adherent K562 erythroid cells (Milon et al. [2001](#page-11-0); Gaither and Eide [2001\)](#page-10-0). The variation in

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cellular site of localisation of hZip1 raises questions about its role in directly mediating cellular Zn accumulation.

That the primary function of Zip1 may not be the direct acquisition of Zn is suggested by the expression of Zip1 in the stromal compartment of the developing mouse gut and relative lack of effect of dietary Zn on Zip1 mRNA levels in the adult mouse intestine (Huang et al. [2006](#page-10-0)) and in human HT-29 colorectal cells (Gurusamy et al. [2011](#page-10-0)). Interestingly, the $h\mathbb{Z}ip1$ gene is located within the epidermal differentiation complex (Lioumi et al. [1999\)](#page-10-0) and plays a key role in the differentiation of human bone cells (Khadeer et al. 2005 ; Tang et al. 2006). Mouse $Zipl$ knockout studies suggest that Zip1 has an indirect function in Zn uptake as no adverse effects were seen although mice were unable to adapt to nutritional Zn deficiency (Dufner-Beattie et al. [2006;](#page-10-0) Kambe et al. [2008](#page-10-0)). Thus, Zip1 may be involved in Zn homoeostasis through a regulatory rather than a primary role in cellular Zn uptake.

The purpose of this study was to establish whether hZip1 is localised to the apical plasma membrane where it could facilitate enterocyte Zn uptake and to employ knockdown and over expression experiments to demonstrate a role for hZip1 in cellular Zn uptake.

Materials and methods

Human tissue

Small intestine tissue was collected from gastroendoscopy biopsies from normal subjects. Tissue from normal resting breast was obtained from breast biopsies performed for diagnosis of breast diseases, and skin tissue was obtained from plastic surgery. The tissues were immediately frozen at -80 °C until use. Ethical consent for this study was granted by Deakin University EC 3.2–2000 and by the Royal Children's Hospital EHR 20025 A.

Cell culture

Caco-2 human adenocarcinoma cells were cultured in 75-cm² culture flasks (Ackland et al. 2005) or on EHSmatrix (Sigma, Australia)-coated porous Transwell filters (Costar, Australia) in DMEM medium supplemented with 10 % FBS (CSL, Australia). Frozen cell pellets from human colorectal adenocarcinoma HT29, duodenal adenocarcinoma HuTu80 and human brain neuroblastoma LA-1s cell lines were also used.

Treatment of cells

Cells were treated with 100 μ M ZnCl₂, 100 μ M ZnCl₂ plus 0.2 nM pyrithione, 6 μ M N, N, N, N-tetrakis (2-pyridylmethyl) ethylenediamine (TPEN) (Sigma, Australia), for 48 h in growth media (Wilson et al. [2006](#page-11-0)). Foetal bovine serum (FBS) was incubated with analytical grade Chelex 100 resin 200–400 sodium mesh (Biorad, Australia) to remove zinc by chelation (Wilson et al. [2006](#page-11-0)). The zinc concentration in untreated and treated serum was measured by the atomic absorption spectrophotometry (Varian Spectra AA-800 (Varian, USA) and indicated a reduction in zinc content from 41 μ M in untreated serum to 1.2 μ M in serum incubated with Chelex 100. The levels of copper and iron were unchanged with Chelex treatment. Chelex-treated serum was stored at $4 °C$ and added to the media when required.

hZip1 cDNA, plasmid construction and generation of transfected Caco-2 lines

The GenBank NM_014437 sequence was selected as it contained the complete open reading frame (ORF) of $hZIP1$ gene. A small fragment of the 5' region of hZIP1 ORF was amplified with primers, ZIP1-C (GGTCTGAG AGTCACTGGAGC) and ZIP1-E (GAGCGTCACGTGC AAGGCTG), from a range of cells and tissues.

The $hZIP1$ coding sequence was amplified using the ZIP1-1F (ATAGAATTCATGGGCCTGGGGGAGAGC) and ZIP1-2R (AAATCTCGAGCTA GATTGGATGAA GAGCAG) primers containing *EcoRI* and *XhoI* restriction sites, respectively, and cloned into a pcDNA3 mammalian expression vector (Life Technologies, Australia). The pcDNA-hZIP1 plasmid was grown in and isolated from E. coli. This plasmid was transfected into Caco-2 cells using LipofectAMINE 2000 reagent (Life Technologies, Australia) following the manufacturer's instructions. The cells stably expressing the hZIP1 from CMV promoter were isolated using G418 (500 µg/ml) (Life Technologies, Australia) selection.

Plasmids containing a myc-tagged sequence, hZIP1 myc, were constructed through similar procedures; however, the hZIP1 sequence was amplified using two primer sets to allow for insertion of the c-myc sequence into the region between transmembrane domains II and III and was predicted by TOPCONS (<http://topcons.cbr.su.se/>) software to face the extracellular compartment. Primers ZIP1 myc1F (ATAGAATTCATGGGCCTGGGGGAGAGC) containing EcoRI site and ZIP1-myc1R (AATACTAGTC AGGTCCTCTTCAGAGATAAGTTTTTGTTCAGCCAG GTAGTCAGGCA), consisting of 20 nucleotides of complementary sequence to hZIP1 cDNA, and encoding the c-myc gene fragment with BamHI restriction site, were used to amplify first fragment of hZIP1-myc construct. The second fragment was amplified using primers containing the BamHI site and ZIP1-myc2F (ATAACTA GTGCCATAGATGAGGCCCTGGCA) and ZIP1-myc2R

(AAATCTCGAGCTAGATTGGATGAAGAGCAG) containing XhoI site. Both fragments were digested using EcoRI and BamHI or BamHI and XhoI restriction enzymes, respectively, and ligated together using T4 DNA ligase system (Roche, Australia) according to the manufacturer's instructions, then cloned into a pcDNA3 vector. Transfection procedures and clone selection were performed as for hZIP1 construct.

siRNA: Two sets of oligonucleotides (21-m) were designated, matching the hZIP1 ORF sequence: siZIP1-1a (AAGGCTCAGCTTCCCGCCAGACCTGTCTC), siZIP1- 1b (AATCTGGCGGGAAGCTGAGCCCCTGTCTC), siZIP1-5a (AACCCCCTCAGCCTTGCGTGCCCTGTCTC) and siZIP1-5b (AATCCTGACCCTCTCCCTGTTCCT GTCTC). Additionally, control siRNA oligonucleotides siCON-1a (AATGCATGTGTCATCGTGATACCTGTC TC) and siCON-1b (AATATGACGATGACACATGCAC CTGTCTC) lacking significant homology to any human transcript were designed. All oligonucleotides were used for in vitro transcription with T7 RNA polymerase following the manufacturer's protocol 'Silencer siRNA construction kit' (Ambion, Australia). Caco-2 cells were transiently transfected with either 50 nM hZIP1 siRNA or 50 nM control siRNA using DMRIE-C reagent and Opti-MEM media (Invitrogen Life Technologies, Australia) following the manufacturer's protocol. Cells were used for experiments 56 h after transfection.

⁶⁵Zn accumulation

The pcDNA-hZIP1, vector control and siRNA-transfected Caco-2 cell were grown to confluency in 6-well plates (Costar, Australia). Growth media were removed from cells and replaced with pre-warmed media supplemented with 1.5 μ Ci/ml of carrier-free ⁶⁵Zn (Perkin Elmer, Australia), and then, cells were incubated for 60 min. The total zinc concentration of the media was $4.1 \mu M$. The cell-associated radioactivity was measured with Minaxi Auto Gamma counter, model 5530 (Packard Instrument Company, Illinois, USA). Zn accumulation was normalised to the protein concentration of lysed cells, estimated using Bio-Rad D_c protein assay (Bio-Rad, Australia).

Real-time PCR

Real-time PCR was performed as previously described (Michalczyk et al. [2003\)](#page-10-0). The primers ZIP1realA (GCCC TGAGCCTAGTAAGCTGTT) and ZIP1realB (TCATCTA TGGCAGCCAGG TAGTC) were used to amplify hZIP1 fragments. Primers 18s-realA (CGGCTACCACATCCAA GGAA) and 18s-realB (GCTGGAATTACCGCGGCT) designed to the 18s ribosomal sequence, an internal control, were run in parallel reactions. A DNA-free kit (Ambion, Australia) was applied to remove genomic DNA. $2^{-\Delta\Delta CT}$ and relative mRNA quantity calculations and statistical analysis (t test) were performed on all reactions.

Antibodies

The sequence encoding the N-terminus of hZIP1 transporter was used to raise antibodies. Briefly, the ZIP1-myc1F forward primer and GST-R reverse primer (AATCTCGAG AACTTCACCTCCAGCCCCAC) were utilised to amplify the hZIP1 fragment with restriction sites for EcoRI and SpeI correspondingly. The PCR product was cloned into pGEX-2T (Amersham Pharmacia, Sweden) in frame with the carboxy terminus of the bacterial glutathione-S-transferase (GST) gene. The GST-hZIP1 fusion protein was expressed in E. coli according to the manufacturer's protocol. The GST-hZIP1 fusion protein was used to raise antibodies (hZip1-FP) to the hZIP1 transporter in sheep (Institute of Medical and Veterinary Science, Adelaide, South Australia).

The following commercially available antibodies and reagents were used: monoclonal anti-myc antibody (Roche, Melbourne, Australia), monoclonal anti-human transferrin receptor antibody and phalloidin-FITC conjugant (Sigma-Aldrich, Australia), monoclonal anti-PDI (protein disulphide isomerase) antibody (Stressgene, Canada), monoclonal anti-villin (Immunotech, Slovak Republic), monoclonal anti-p230 (Transduction Laboratories, USA) and monoclonal anti-E-cadherin (Chemicon, Australia).

Immunofluorescence

Adherent cultured cells were grown on 10-mm-diameter glass cover slips and processed for immunostaining as previously described (Michalczyk et al. [2003\)](#page-10-0). Briefly, selected primary antibodies were diluted in 1 % BSA in PBS and applied to cells overnight at 4° C. Control cells were incubated with ZIP1-FP fusion protein overnight. After 3 PBS washes, a secondary antibody AlexaFluor 488 donkey antisheep IgG or donkey anti-mouse IgG (Molecular Probes, USA), 1 in 2000 dilution was applied for 2 h at room temperature. Additional Cy-5 donkey anti-sheep or donkey antimouse secondary antibodies (Chemicon, Australia) were added to cells when double label was required. To label only the surface pool of hZIP1-myc, hZIP1 and transferrin receptor, the Triton X-100 permeabilisation step was omitted and cells were immunolabelled as described above. Plasma membrane binding and internalisation of anti-hZIP1, antimyc and anti-transferrin receptor antibodies were detected when live cells were incubated with $5 \mu g/ml$ antibodies for 1 h. After that, cells were fixed, permeabilised and processed as described previously. Cells were viewed using a Leica TCS SP2 AOBS laser confocal microscope (Leica,

Australia), with PlanApo 100x oil objective at RT. Images were viewed and captured on a HP workstation with Leica Microsystems TCS SP2 software (Leica, Australia). Image overlay, brightness, contrast and colour balance adjustments were done using Photo Paint software version 6.0.

Statistics

Data are presented as mean \pm SD. Statistical analyses were carried out using a *t* test with unequal variance.

Results

hZip1 was expressed endogenously in the human small intestine and in cultured human gut cell lines

Expression of hZip1 mRNA was detected in extracts from human small intestine and in extracts from cultured gut cells. For comparison, we also investigated hZip1 in other human tissues including breast and skin, and in cultured human skin fibroblasts and lymphoblasts, using RT-PCR. A band of size 341 bp was detected in all samples (Fig. 1).

hZip1 protein had a perinuclear distribution in nondifferentiated Caco-2 cells that was not influenced by extracellular zinc conditions

The localisation of endogenous hZip1 protein in Caco-2 cells was determined. An antigen/antibody competition

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20

Fig. 1 hZip1 gene product was detected in a wide range of human tissues and cell lines. \bf{a} A band (341 bp) representing the 5' region of the hZIP1 ORF was found in all tissues and cells tested using RT-PCR. PMC42 (human breast adenocarcinoma) cells (lane 2), Caco-2 (human colon adenocarcinoma) cells (lane 3), HT29 (human colon adenocarcinoma) cells (lane 4), HuTu80 (human duodenum adenocarcinoma) cells (lane 5), LA-1 s (human neuroblastoma) cells (lane 6), fibroblasts 1 (lane 7), fibroblasts 2 (lane 8), fibroblasts 3 (lane 9), fibroblasts 4 (lane 10); fibroblasts 5 (lane 11), lymphoblasts 1 (lane 12), lymphoblasts 2 (lane 13), lymphoblasts 3 (lane 14), small intestine biopsy 1 (lane 15), small intestine biopsy 2 (lane 16), resting breast tissue 1 (lane 17); resting breast tissue 2 (lane 18), finger skin tissue (lane 19)

control showed no staining (Fig. [2](#page-4-0)A, a). A pattern of fine cytoplasmic granules concentrated in the perinuclear region was detected in Caco-2 cells probed with hZip antibody (Fig. [2A](#page-4-0), b). Similar patterns of distribution of hZip1 were seen in Caco-2 cells treated for 48 h with Zn (Fig. [2](#page-4-0)A, c) and Zn with pyrithione (Fig. [2](#page-4-0)A, d) or subjected to zincdeficient conditions including TPEN (Fig. [2](#page-4-0)A, e) and Chelex-treated serum (Fig. [2](#page-4-0)A, f). The XZ sections confirmed the cytoplasmic localisation of hZip1 in untreated cells (Fig. [2](#page-4-0)A, b') and cells exposed to Zn (Fig. [2A](#page-4-0), c'), Zn with pyrithione (Fig. [2](#page-4-0)A, d'), TPEN (Fig. [2](#page-4-0)A, e') and grown in Chelex-treated media (Fig. [2](#page-4-0)A, f').

In Caco-2 cells, hZip1 mRNA decreased in response to Zn treatment and increased in response to Zn depletion

The responsiveness of $hZip1$ mRNA to various Zn treatments was tested. Zinc concentrations of 100 μ M Zn, 100 μ M Zn and 0.2 nM pyrithione, 6 μ M TPEN and Chelex-100-treated serum were added to Caco-2 cells for 48 h. Additionally, Caco-2 cells were grown on Transwell filters coated with EHS matrix to induce cell differentiation. A significant reduction in hZip1 mRNA levels in response to zinc and zinc with ionophore treatment (2.7 fold with Zn, 3.2-fold with Zn and ionophore) was found, while zinc-deficient condition significantly increased $hZip1$ transcript abundance (6.4-fold with TPEN, 5.8-fold with Chelex-treated serum) (Fig. [2](#page-4-0)B). Differentiated Caco-2 demonstrated a slight (2.1-fold) but significant increase in quantity of $hZip1$ mRNA (Fig. [2](#page-4-0)B).

In non-differentiated Caco-2 cells, hZip1 was partially localised to the ER compartment

To identify the subcellular compartment in which endogenous hZip1 was localised, antibodies directed against proteins of known localisation within the cell were used together with the hZip1 antibody in Caco-2 cells (Fig. [3a](#page-5-0), e, i, m, green). Nuclei were stained with ethidium bromide and pseudo-coloured in blue (Fig. [3](#page-5-0)c, g, k, o). hZip1 protein in Caco-2 cells showed a cytoplasmic granular label (gr) with brighter perinuclear staining (pr) in some cells (Fig. [3a](#page-5-0), e, i, m) in contrast to the junctional label seen with E-cadherin (Fig. [3](#page-5-0)b) with no colocalisation between Zip1 and E-cadherin (Fig. [3d](#page-5-0)). The transferrin receptor showed a punctate cytoplasmic pattern with some larger granules visible (lg) (Fig. [3f](#page-5-0)), and the overlay (Fig. [3](#page-5-0)h) showed no colocalisation between hZip1 and transferrin receptor labels. The immunostaining of the p230 protein (Fig. [3](#page-5-0)j), which is localised to the trans-Golgi compartment, was distinctively different from that of hZip1

Fig. 2 Subcellular localisation of hZip1 protein in Caco-2 cells was not altered by zinc levels but differences in the expression of endogenous $hZip1$ mRNA were detected in response to changes in zinc treatment and differentiation. A Caco-2 cells were grown in control medium (b), Zn-supplemented (c and d) or Zn-deficient (e and f) media for 48 h and incubated with hZip1 Ab. Cells were stained with hZip1 antibody with fourfold excess of antigen to test the Ab specificity (a). Untreated medium (b), medium supplemented with 100 μ M zinc (c), medium with 100 μ M zinc and 0.2 nM pyrithione (d) zinc-deficient medium with 6 μ M TPEN (e) and also Chelex-

seen in the overlay (Fig. [3l](#page-5-0)). The immunostaining of PDI (Fig. [3](#page-5-0)n), a resident protein of the ER, showed a cytoplasmic granular label (gr) with brighter perinuclear staining in some cells (pr). The overlay with

treated medium (f). The XZ sections (b'–f'). B Real-time PCR analysis showed down-regulation on hZip1 mRNA in Caco-2 cells exposed to 100 μ M zinc (grey bar) and 100 μ M zinc with 0.2 nM pyrithione (blue bar), while 6 µM TPEN treatment (light green bar) and incubation with Chelex-treated media (dark green bar) resulted in increased levels of hZIP1 transcript. In differentiated Caco-2 upregulation, the hZip1 levels were detected (dark grey bar) compared to the control (solid white bar). The bars represent the mean $(\pm SD)$ of 3 independent experiments. Stars indicate the statistically significant differences ($p < 0.05$, t test)

hZip1 protein (Fig. [3](#page-5-0)p) demonstrated partial colocalisation (yellow colour) between hZip1 and PDI labels, especially in perinuclear region (pr) and in granules located close to this region (pg).

Fig. 3 hZip1 was partially colocalised to the ER in non-differentiated Caco-2 cells. hZip1 protein showed cytoplasmic granular (gr) and perinuclear staining in some cells (pr) (a, e, i and m). E-cadherin showed a punctate cell junction pattern (b) with no colocalisation between hZip1 and E-cadherin (d). Transferrin receptor was detected as a punctate cytoplasmic pattern (f). The overlay showed no colocalisation between hZip1 and transferrin receptor (h). trans-Golgi

In differentiated Caco-2 cells, hZip1 was localised beneath the microvilli

To investigate the localisation of hZip1 in differentiated gut epithelia that more closely resemble enterocytes, Caco-2 cells were grown on EHS-coated Transwell filters for 3 weeks to induce differentiation and then stained with the hZip1 antibody and cellular markers. E-cadherin label was detected at the cell junctions (Fig. [4](#page-6-0)g), while hZip1 had a granular cytoplasmic distribution (Fig. [4d](#page-6-0)). There was no significant colocalisation between E-cadherin and hZip1 proteins within differentiated Caco-2 cells in XY or XZ sections (Fig. [4j](#page-6-0), j'). F-actin was expressed mainly at the vicinity of the apical region of the cells (Fig. [4](#page-6-0)h) and showed some overlap with

marker p230 demonstrated perinuclear label (j) with no colocalisation with hZip1 (I). PDI protein localisation was similar to hZip1 (n) with partial colocalisation (yellow colour) between these proteins was identified in perinuclear region (pr) and in granules located close to this region (pg) . Cell nuclei were stained with ethidium bromide (c, g) , k and o)

hZip1 (Fig. [4k](#page-6-0), k'). Villin was detected only at the apical surface of Caco-2 cells marking the position of microvilli (Fig. [4i](#page-6-0)). The hZip1 label appeared to overlap with villin (Fig. [4l](#page-6-0)) but the XZ section showed that hZip1 was present beneath the microvilli (Fig. [4](#page-6-0)l'). Our hZip1 immunostaining shows that it is possible that in differentiated cells, some Zip1 protein remains within the Golgi network.

Internalisation studies indicated that hZip1-myc protein was not present at the plasma membrane of Caco-2 cells

To confirm that hZip1 was not located at the plasma membrane but adjacent to the apical surface in

differentiated cells, a myc-tag was incorporated into the predicted extracellular loop of hZip1 protein and the construct introduced into Caco-2 cells. Caco-2 cells expressing the hZip1-myc were used for trafficking experiments to enable us to determine whether the hZip1 protein resided at the plasma membrane or alternatively, near the plasma membrane. Western blot analysis of extracts from Caco-2 cells transfected with pcDNA3-hZip1-myc showed a band of approximately 35 kDa, the expected size for hZIP1-myc protein, with anti-myc antibodies (Fig. [5A](#page-7-0) lane 2) but not in the vector control (Fig. [5](#page-7-0)A lane 1). In the permeabilised Caco-2 cells, hZip1-myc showed fine-granular, cytoplasmic label (Fig. [5B](#page-7-0)c) not seen in cells transfected with vector alone (Fig. [5B](#page-7-0), a). In Caco-2 cells, no staining was found on the surface of the non-permeabilised cells (Fig. [5](#page-7-0)B, d) proving that hZip1 was not localised to the plasma membrane.

To test the possibility that hZip1 may be located to the plasma membrane under conditions of zinc deficiency and subsequently endocytosed by cells, the Caco-2-hZip1-myc cells were incubated in either growth medium or 6μ M TPENsupplemented growth medium in the presence of $5 \mu g/ml$ anti-myc antibodies for 60 min and then processed for immunofluorescence. Antibody $(5 \mu g/ml)$ to the extracellular region of the transferrin receptor, a protein which traffics between plasma membrane and early endosomes, was used as internalisation control for the Caco-2-hZip1-myc cells. The hZip1-myc protein was not detected on the surface or internalised by Caco-2 cells grown in either zinc-sufficient or zincdeficient media (Fig. [5](#page-7-0)C, a, b). However, the transferrin receptor showed strong granular cytoplasmic staining including plasma membrane label under these two conditions (Fig. [5C](#page-7-0), c and d). These results do not support the idea of constitutive or zinc-induced cycling of hZip1 to the plasma membrane.

Fig. 5 The hZip1-myc protein was not detected at the plasma membrane of Caco-2 cells, and hZip1-myc-tagged protein was not internalised by the cells. A Western blot analysis of the Caco-2 cell extracts probed with anti-myc antibody, with β -actin expression as loading control, representative of 3 independent experiments. Lane 1, Caco-2 cells transfected with pcDNA3 vector control; Lane 2, Caco-2 cells transfected with hZip1-myc construct, showing 35 kDa band of expected size. B immunolocalisation of hZip1 protein in Caco-2 cell line. Permeabilised (a, c) and non-permeabilised (b, d) cells were labelled with anti-myc antibody. No staining in vector control Caco-2 cells (a and b). No label in non-permeabilised Caco-2 (d) cells. Permeabilised Caco-2 cells had a granular cytoplasmic pattern (c). C Live cells were incubated with $5 \mu g/ml$ anti-myc or anti-transferrin receptor antibodies for 1 h. The hZip1-myc Caco-2 cells were either maintained in growth media (a, c) or in media supplemented with 6 μ M TPEN (b, d). Surface binding or internalisation of anti-myc antibody was not detected in cells (a, b) while anti-transferrin receptor antibody was found at the plasma membrane and in the cytoplasm (c,)

Overexpression of hZip1 transporter increased Zn accumulation in non-differentiated Caco-2 cells

To determine the role of hZip1 in zinc transport, the complete Zip1 ORF was cloned into the pcDNA3 mammalian expression vector and stably expressed from a CMV promoter in Caco-2 cells. Real-time PCR was used to quantify hZip1 transcripts using the 18s ribosomal subunit as an internal control. In hZip1 overexpressing Caco-2 cells, the hZip1 levels were significantly increased to three times that of the vector-only cells (Fig. [6](#page-8-0)a). To verify whether overexpression of hZip1 was associated with increased zinc accumulation, cells were exposed to ${}^{65}Zn$ in DMEM medium supplemented with 10 % FBS for 1 h. Accumulation of zinc in Caco-hZip1 cells (208.5 pmol Zn/ mg protein) was significantly increased by twice relative to that of the Caco-2-vector control cells (115 pmol Zn/mg protein) ($p < 0.05$, t test) (Fig. [6](#page-8-0)b).

Knockdown of hZip1 reduces Zn accumulation in non-differentiated Caco-2 cells

A function for the endogenous hZip1 protein in transporting zinc was investigated by assessing the effect of inhibition of hZIP1 expression on ⁶⁵Zn accumulation. Caco-2 cells were transfected with two different siRNAs designed to hybridise with coding region of the hZip1 mRNA and to cause degradation of the transcript by the RNA-induced silencing complex. The $hZipP1$ mRNA level, estimated by real-time PCR, was significantly reduced by 2.2-fold when the cells were treated with 50nM of siRNA hZip1 compared to cells treated with 50 nM randomly selected control siRNAs, with no similarity to any known human mRNA (Fig. [6c](#page-8-0)). The ⁶⁵Zn accumulation in $hZip1$ siRNA-transfected Caco-2 cells was 20 % lower than in control siRNA cells ($p > 0.05$, t test) (Fig. [6d](#page-8-0)).

Discussion

Zip1 is one of the 14 members of the SLC39 putative Zn transporters that are postulated, but not conclusively shown to mediate cellular accumulation of Zn. We detected hZip1 mRNA transcripts in human intestinal tissue and in a cultured human gut line as well as in other human tissues including breast and skin and cultured human skin fibroblasts and lymphoblasts, indicating widespread expression of the human gene that is consistent with the expression in mouse tissues (Dufner-Beattie et al. [2003\)](#page-10-0). An explanation for the presence of multiple members of the SLC39A is not clear but it would seem that the role of these transporters may extend beyond that of a transmembrane Zn transporter.

Fig. 6 Zinc accumulation was increased in Caco-2 cells overexpressing the hZip1 construct and decreased in cells transfected with hZip1 siRNA. a hZip1 mRNA expression levels in Caco-2 cells transfected with hZip1 constructs were higher relative to vector. b Zinc accumulation was increased in Caco-2 cells transfected with hZip1 construct relative to vector-only following incubation in ${}^{65}Zn$. c $hZip1$ mRNA expression levels in Caco-2 cells transfected with hZip1 siRNA was reduced relative to control siRNA. d Zinc accumulation in Caco-2 cells transfected with hZip1 siRNA was reduced compared to control siRNA following incubation in ${}^{65}Zn$. The *bars* represent the mean $(\pm SD)$ of 3 independent experiments. Stars indicate the statistically significant difference ($p < 0.05$, t test)

In cultured human Caco-2 cells, endogenous hZip1 protein had a cytoplasmic, vesicular location that partially overlapped with the endoplasmic reticulum marker PDI, similar to the localisation seen in other studies (Milon et al. [2001;](#page-11-0) Khadeer et al. [2005](#page-10-0)). In contrast, in Caco-2 cells that were differentiated by growth on EHS matrix-coated filters, the Zip1 protein was detected in the vicinity of the apical membrane. This indicates that the change in Zip-1 localisation was associated with the differentiated state. It is consistent with previous studies showing that the extracellular matrix is essential for proper differentiation (Schock and Perrimon [2002\)](#page-11-0), and the induction of genes is required for the formation of differentiated enterocytes (Halbleib et al. [2007;](#page-10-0) Saaf et al. [2007\)](#page-11-0) and for targeting of molecules to the appropriate domain in enterocytes (Schreider et al. [2002](#page-11-0)). Previous studies with Caco-2 cells have shown that hZip1 expression is influenced by the differentiation state of the cells (Zemann et al. [2010\)](#page-11-0). Our data have parallels with previous findings on the role of frataxin in iron transport in differentiated Caco-2 cells. The authors reported that the frataxin localisation changed during Caco-2 cell differentiation, which was attributed to its role in iron intestinal uptake in differentiated cells. It was concluded that differentiated enterocytes are specialised for absorption and transport of iron and differ from

the undifferentiated precursors in their expression of iron uptake and iron transport proteins (Acquaviva et al. [2005](#page-10-0)). The change in localisation of hZip1 in Caco-2 cells could be linked to zinc fluxes, in relation to changes in enterocyte metal absorption following their differentiation.

Colocalisation studies of differentiated Caco-2 cells indicated that although hZip1 was apically located, it did not reside at the plasma membrane but was beneath the microvilli marker, villin. Our functional data indicate no association of the hZip1 protein with the apical plasma membrane. This was apparent from experiments in Caco-2 cells transfected with a myc epitope predicted to be placed in extracellular region (TOPCONS topology software) that showed no hZip1 staining in non-permeabilised or live cells that had been incubated with an anti-myc antibody. Furthermore, endogenously expressed hZip1 protein was not detected at the cell surface with the antibody to the extracellular N-terminus of hZip1 protein, even if the cells were grown in a zinc-deficient environment. In contrast, under the same experimental conditions, the transferrin receptor was detected on the cell surface from where it was internalised into cytoplasmic vesicles.

We found that the localisation of the hZip1 protein was not altered in response to changes in extracellular zinc levels. This is consistent with a similar study on Caco-2 cells (Shen et al. [2008\)](#page-11-0) but in contrast to the intracellular trafficking of hZip1 protein reported in transfected human embryonic kidney cells (Wang et al. [2004\)](#page-11-0) and in CHO cells (Huang and Kirschke [2007\)](#page-10-0). Thus, hZip1 Zn responsiveness appears to be cell type specific.

Although we did not detect the hZip1 protein at the plasma membrane of differentiated Caco-2 cells, where it would be predicted to be if it was involved in Zn uptake, we demonstrated a Zn transport role for the endogenous protein by the 20 % reduction in zinc accumulation in hZip1 knockdown cells. When hZip1 was overexpressed in Caco-2 cells, 65Zn accumulation was increased by twofold. This is similar to other studies in human K562 erythroleukemia cells (Gaither and Eide [2001](#page-10-0)) and PC-3 prostate cells (Franklin et al. [2003\)](#page-10-0) where overexpression of Zip1 induced Zn accumulation and the rate of Zn uptake. Thus, our study as well as those of others unequivocally demonstrates that Zip1 plays a role in Zn accumulation.

We tested the Zn responsiveness of $hZip1$ in differentiated Caco-2 cells by exposing them to excess $Zn(100 \mu M)$ Zn and 100 μ M Zn + zinc ionophore) and to Zn depletion (Zn chelator and Chelex-100-treated serum). Significant decreases in $hZip1$ mRNA were seen in response to excess Zn, while Zn deficiency induced increase in the levels of $hZip1$ mRNA. It should be noted that in the mouse, intestine dietary levels of Zn did not affect the abundance Zip1 mRNA or protein (Huang et al. [2006](#page-10-0); Dufner-Beattie et al. [2003\)](#page-10-0), contrary to the human gut, where the response to dietary Zn differs from the rodent model (Cragg et al. [2005\)](#page-10-0).

To account for our findings and explain the function of hZip1 in human enterocytes, we developed a hypothetical model. Zn is taken up by ion channels such as the K^+/Zn^{2+} antiporter (Ackland and McArdle 1990, 1996) or the Na⁺/ Zn^{2+} counter transporter (Ohana et al. [2004\)](#page-11-0) or transporters such as Zip4 (Wang et al. [2002\)](#page-11-0) and the multi-ion importer DCT1/DMT1/Nramp2 (Gunshin et al. [1997\)](#page-10-0); however, the

affinity of DMT1 for zinc is low (Yamaji et al. [2001](#page-11-0)). Imported zinc is delivered to zinc-dependant proteins and sequestered into vesicles by SLC30/CDF transporters including ZnT2 (Palmiter et al. [1996a](#page-11-0)), ZnT3 (Palmiter et al. [1996b](#page-11-0)) and ZnT4 (Michalczyk et al. [2002](#page-10-0); Murgia et al. [1999\)](#page-11-0). The zinc-containing vesicles or zincosomes may be utilised for temporary metal storage or directed into an exocytotic pathway. Zinc is released from vesicular storage by SLC39A transporters in the vesicular membrane in response to decreases in cytoplasmic Zn levels.

In the cytoplasmic loop between transmembrane domains III and IV, two histidines conserved within the SLC39A family are potential Zn-binding regions required for Zn transport (Milon et al. [2006\)](#page-11-0) and may act as a zinc sensor (Fig. 7) similar to the zinc-binding region in the yeast ZAP1 transcription regulator (Bird et al. [2000](#page-10-0)). If intracellular levels of Zn increase, more Zn is bound by hZip1 and this prevents release of stored vesicular Zn into the cytoplasm. It is also possible that hZip1, activated by zinc binding, induces additional mechanisms controlling the vesicular zinc storage, therefore contributing to intracellular zinc homoeostasis. When the cytoplasmic Zn levels fall, less Zn is bound to hZip1, resulting in a translocation of the metal from vesicles to the cytoplasm (Fig. 7). The feasibility of this hypothetical model is indicated by examples of similar systems involving Zn sensors, including the store-operated calcium channel (SOCC) where Zn inhibits calcium influx (Gore et al. [2004\)](#page-10-0) and Yiip, a member of the SLC30 family which effluxes Zn in response to the activation of the cytoplasmic domain (Lu and Fu [2007;](#page-10-0) Nies [2007](#page-11-0)).

Based on our experimental data, we propose a hypothesis that would explain the observations of higher intracellular Zn concentrations in cells overexpressing hZip1 compared to control cells. In hZip1 overexpressing cells, there is an increased number of hZip1 molecules present on the vesicular membranes that would bind zinc by the

Fig. 7 A model for hZIP1 as intracellular zinc sensor in Caco-2 cells. a Zinc is taken up from the extracellular environment by transporter X. To maintain intracellular zinc homoeostasis, endogenous hZIP1 regulates zinc release from zinc-containing vacuoles (zicosomes) into

the cytoplasm in basal zinc and zinc-deficient conditions by utilising 2 cytoplasmic histidines as zinc concentration sensors. This transport is ceased following an increase in extracellular/cytoplasmic zinc levels as the hZIP1 sensors are saturated

histidine/sensor region. The Zn in the cytoplasm binds to hZip-1 until saturation of histidines on additional exogenous hZip1 is achieved and Zn efflux from the vesicles is prevented. In addition, hZip1 may also induce activation of additional mechanisms to facilitate zinc transport into vesicles that could account for the overall increase in zinc accumulation in the hZip1 overexpressing cells. The silencing of the *hZip1* gene produces the opposite effect, where fewer transporters are available to bind Zn; therefore, equilibrium is established at a lower Zn concentration. As a regulator of zinc fluxes, hZip1 may also contribute to zinc signalling within the cells in addition to its role in calcium regulation (Maret 2001, 2011; Hershfinkel et al. 2001).

In conclusion, the hZip1 protein was not a resident of the apical plasma membrane and does not directly facilitate Zn accumulation in differentiated Caco-2 cells, although it is zinc-responsive. It is possible that hZip1 may function as a Zn sensor that has a regulatory role in zinc homoeostasis in human gut enterocytes, similar to that of SOCC or Yiip.

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