ZIP8 is an iron and zinc transporter whose cell-surface expression is upregulated by cellular iron loading*

Chia-Yu Wang1, Supak Jenkitkasemwong1, Stephanie Duarte1, Brian K. Sparkman2, Ali Shawki2,3, Bryan Mackenzie1, and Mitchell D. Knutson1

1From the Food Science and Human Nutrition Department, University of Florida, Gainesville, Florida 32611

2Department of Molecular & Cellular Physiology, and 3Systems Biology & Physiology Program, University of Cincinnati College of Medicine, Cincinnati, Ohio, 45267

*Running title: ZIP8 is an iron-regulated iron and zinc transporter

To whom correspondence should be addressed: Mitchell D. Knutson, PO Box 110370, University of Florida, Gainesville, FL 32611; Fax: 352-392-9467; E-mail: mknutson@ufl.edu

**Background:** Previous studies have identified the transmembrane protein ZIP8 (ZRT/IRT-like Protein 8) as a zinc transporter.

**Results:** ZIP8 can transport iron in addition to zinc and is upregulated by iron loading.

**Conclusion:** ZIP8 represents the third mammalian transmembrane iron-import protein to be identified.

**Significance:** ZIP8 may play a role in iron metabolism.

**SUMMARY**

ZIP8 (SLC39A8) belongs to the ZIP family of metal-ion transporters. Among the ZIP proteins, ZIP8 is most closely related to ZIP14, which can transport iron, zinc, manganese, and cadmium. Here we investigated the iron transport ability of ZIP8, its subcellular localization, pH dependence, and regulation by iron. Transfection of HEK 293T cells with ZIP8 cDNA enhanced the uptake of 59Fe and 65Zn by 200% and 40%, respectively, compared with controls. Excess iron inhibited the uptake of zinc and vice versa. In RNA-injected *Xenopus* oocytes, ZIP8-mediated 55Fe2+ transport was saturable (K0.5 of ≈ 0.7 μM) and inhibited by zinc. ZIP8 also mediated the uptake of 109Cd2+, 57Co2+, 65Zn2+ > 54Mn2+, but not 64Cu (I or II). By using immunofluorescence analysis, we found that ZIP8 expressed in HEK 293T cells localized to the plasma membrane and partially in early endosomes. Iron loading increased total and cell-surface levels of ZIP8 in H4IIE rat hepatoma cells. We also determined by using site-directed mutagenesis that asparagine residues 40, 88, and 96 of rat ZIP8 are glycosylated and that N-glycosylation is not required for iron or zinc transport. Analysis of 20 different human tissues revealed abundant ZIP8 expression in lung and placenta, and that its expression profile differs markedly from ZIP14, suggesting nonredundant functions. Suppression of endogenous ZIP8 expression in BeWo cells, a placental cell line, reduced iron uptake by ~40%, suggesting that ZIP8 participates in placental iron transport. Collectively, these data identify ZIP8 as an iron transport protein that may function in iron metabolism.

The mechanisms by which iron is taken up by mammalian cells are poorly understood except for those in just two cell types: the enterocyte and the erythroid precursor cell. Enterocytes take up dietary non-heme iron via the transmembrane protein divalent metal-ion transporter-1 (DMT1) located on the apical membrane (1,2). In erythroid precursor cells, DMT1 transports iron from transferrin-containing endosomes to the cytosol (3). Accordingly, Dmt1 (Slc11a2) knockout mice display iron-deficient erythropoiesis and anemia and intestine-specific Dmt1 knockout mice become progressively anemic after birth, likely
because of an inability to absorb dietary iron (1). Interestingly, in Dmt1 knockout mice, most other tissues developed normally, including the liver, which displayed elevated concentrations of non-heme iron at post-natal day 3 (1). These observations demonstrate that alternative iron uptake pathways must exist. One such pathway may involve the plasma membrane protein ZIP14. Originally identified as a zinc transporter (4), ZIP14 was subsequently shown to transport iron in addition to zinc (5). In contrast to DMT1, which transports iron optimally at pH 5.2-5.5 (6-8), ZIP14 exhibits maximal iron transport at pH 7.5 (5,9), making it well-suited for iron uptake from the plasma, such as from non-transferrin-bound iron (NTBI) during iron overload. Endogenous ZIP14 in HepG2 hepatoma cells has additionally been detected in transferrin-containing endosomes, where it mediates, at least in part, the assimilation of iron from transferrin (10).

The ability of a ZIP family protein such as ZIP14 to transport iron is not without precedent. Indeed, the founding member of the ZIP family, IRT1 (iron-regulated transporter 1) in Arabidopsis thaliana, was identified as an iron transporter (11). As demonstrated by knockout studies, IRT1 is responsible for iron uptake from the soil into the root (12). Arabidopsis IRT2, a close homologue of IRT1, was also found to be a high-affinity iron transporter (13). Additional ZIP family members that transport iron include ZupT in Escherichia coli (14) and LIT1 in Leishmania amazonensis (15).

Among the fourteen mammalian ZIP family members, ZIP14 is most closely related to ZIP8 (16). Mouse ZIP14 and ZIP8 are similar in length (489 vs. 462 amino acids), ~50% of their amino acids are identical, and they each contain a long extracellular N-terminal region with multiple potential glycosylation sites. Notably, ZIP14 and ZIP8 are 90% identical (19/21 amino acids) in putative membrane spanners IV and V, which have been proposed to form a metal translocation pore (17). Given the high degree of similarity between the two proteins, we considered the possibility that ZIP8, like ZIP14, would be able to transport iron. ZIP8 has been shown to transport zinc, cadmium, and manganese (18,19), but measurement of iron uptake has not been reported. Therefore, the primary objective of the present study was to investigate the iron transport ability of ZIP8. We also assessed the regulation of ZIP8 by iron, its subcellular localization, and its expression levels in various human tissues. In addition, we determined which of the potential glycosylation sites of ZIP8 are glycosylated and whether glycosylation is required for metal transport.

EXPERIMENTAL PROCEDURES

Cell culture—HEK 293T and H4IIE rat hepatoma cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM; Mediatech). BeWo cells were maintained in F-12K medium with L-glutamine (ATCC). All media were supplemented with 10% (v/v) fetal bovine serum (FBS; Atlanta Biologicals), 100 units/ml penicillin, and 100 µg/ml streptomycin. Cells were maintained at 37 ºC in 5 % CO₂.

Expression of ZIP8 and measurement of iron and zinc uptake—HEK 293T cells were transiently transfected with rat ZIP8 (GenBank accession number BC089844) or empty vector pExpress–1 (Open Biosystems) for 48 h (Fugene HD, Roche). Prior to uptake, cells were washed twice with serum-free medium (SFM) and incubated for 1 h in SFM containing 2% (w/v) BSA to deplete cells of transferrin and to block the nonspecific binding at 37 ºC. For uptake, cells were incubated with 2 µM ⁵⁹Fe-ferric citrate in SFM in the presence of 1 mM L-ascorbic acid for 2 h at 37 ºC with or without a 10-fold molar excess of zinc, followed by 3 washes of iron chelator solution (1 mM bathophenanthroline sulfonate and 1 mM diethylenetriaminepentaacetic acid) to remove any surface-bound iron. Cells were lysed in buffer containing 0.2 N NaOH and 0.2% (w/v) SDS. Radioactivity was determined by gamma counting and protein concentration was determined colorimetrically by using the RC DC protein assay (Bio-rad).

Expression of ZIP8 and ZIP14 in Xenopus oocytes—We performed laparotomy and ovariection on adult female Xenopus laevis frogs (Nasco) under 3-aminoethylbenzoate methanesulfonate anesthesia (0.1% w/v in 1:1 water/ice, by immersion) following a protocol approved by the University of Cincinnati Institutional Animal Care and Use Committee.
Ovarian tissue was isolated and treated with collagenase A (Roche Diagnostics), and oocytes were isolated and stored at 17 °C in modified Barth’s medium as described (20).

ZIP8 cDNAs for rat (GenBank accession number BC089844) and mouse (GenBank accession number BC006731) were subcloned into the pOX(+) oocyte expression vector between KpnI and NotI sites and under the SP6 promoter. ZIP8 cDNA–pOX constructs were linearized by using NotI and RNA was synthesized in vitro by using the mMESSAGE mMACHINE/SP6 RNA polymerase transcription kit (Applied Biosystems/Ambion) according to the manufacturer’s protocol. We prepared mouse ZIP14 RNA from ZIP14–pOX(+) as described (2). Defolliculate stage V–VI oocytes were injected with 50 ng of RNA and incubated for 5 days (ZIP14) or 8–9 days (ZIP8) before being used in functional assays.

Functional assays in oocytes—We measured radiotracer metal-ion uptake in oocytes as described (2) or as follows. Oocytes were incubated at room temperature (23 °C) in transport medium containing 100 mM NaCl, 2 mM KCl, 2 mM CaCl₂, and 1 mM MgCl₂, and buffered using 0–5 mM 2-(N-morpholino)ethanesulfonic acid and 0–5 mM N’N’-diethylpiperazine (GFS Chemicals, Columbus, OH) to obtain pH 7.5 or as otherwise indicated in Fig. 2C. Media containing Fe²⁺ or Cu²⁺ also contained 1 mM L-ascorbic acid; media containing Cu¹⁺ or Cu²⁺ also contained 100 μM L-histidine. We used ⁵⁵Fe (added as FeCl₃) at final specific activity 220–340 MBq.mg⁻¹, ⁵⁵Mn (added as MnCl₂) at final specific activity 24 MBq.mg⁻¹, and ⁵⁷Co (added as CoCl₂) at final specific activity 0.4–2.9 GBq.mg⁻¹, each obtained from Perkin-Elmer Life Science Products (Boston, MA); ¹⁰⁹Cd (added as CdCl₂) at final specific activity 29 MBq.mg⁻¹ and ⁶⁵Zn (added as ZnCl₂) at final specific activity 63 MBq.mg⁻¹, obtained from the National Laboratory (Oak Ridge, TN); and ⁶⁴Cu (added as CuCl₂) at final specific activity 2.0 GBq.mg⁻¹, obtained from Washington University–St Louis (St Louis, MO). Radiotracer metal-ion uptake was measured over 10 min, i.e. well within the linear phase of uptake in oocytes expressing ZIP14 (2).

Concentration-dependence data were fit by a modified Michaelis–Menten function (Equation 1) for which \( V \) is the velocity (uptake) of substrate \( S \) (⁵⁵Fe²⁺ or ⁵⁷Co²⁺), \( V_{max} \) the derived maximum velocity, \( S \) the concentration of substrate \( S \), and \( K_m \) the substrate concentration at which velocity was half-maximal.

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V = \frac{V_{max}S}{K_m + S}
\]  

Construction of HA-tagged ZIP8—A vector encoding rat ZIP8 with a C-terminal hemagglutinin (HA) tag (YPYDVPDYA) was constructed by using inverse PCR mutagenesis (IPCRM) (21). Briefly, full-length rZIP8 cDNA in pExpress–1 was used as the template with the following two ⁵’-phosphorylated oligonucleotide primers: 5’-GTCGCTGTATTACCGCTAG-CTGGGAGTATGACGTCAGCGCGG-3’ and 5’-GTCGTAAGGGTACTGCAATTCGATGT-CCCTGCGTACAAGG-3’ for the IPCRM reaction. The PCR product was purified by using the QIAquick PCR Purification Kit (Qiagen) and subsequently digested with DpnI to eliminate template DNA. The digested PCR product was ligated using T4 DNA ligase (New England Biolabs) and transformed into bacteria to obtain rZIP8-HA plasmid.

Subcellular localization of ZIP8—HEK 293T cells were grown on poly-L-lysine-coated coverslips and transfected with rat ZIP8-HA. Forty-eight h later, cells were washed twice with PBS buffer containing 1 mM MgCl₂ and 0.1 mM CaCl₂ and fixed with 2% (w/v) paraformaldehyde (PFA) for 15 min at room temperature (RT). Excess PFA was quenched by adding 50 mM NH₄Cl for 10 min, followed by 3 washes of PBS. To permeabilize cells, cells were incubated with 0.1% (w/v) saponin for 10 min and then washed 3 times with PBS before blocking in 1% (w/v) BSA for 30 min. ZIP8-HA was detected by using mouse anti-HA primary antibody (1 µg/mL; Roche) and rabbit anti-mouse IgG Alexa Fluor 594 secondary antibody (1:100; Invitrogen). To confirm permeabilization of cells, the lysosomal marker LAMP-1 (lysosome associated macrophage protein 1) was detected by using goat anti-LAMP-1 primary antibody (1:1000; Santa Cruz Biotechnology) and donkey anti-goat secondary
antibody (1:100; Invitrogen). The endosomal marker, early endosome antigen 1 (EEA1) was detected by using goat anti-EEA1 primary antibody (1:1000; Santa Cruz Biotechnology) and the donkey anti-goat IgG Alexa Fluor 488 secondary antibody (1:100; Invitrogen). To stain nuclei, cells were incubated with 10 µg/mL 4′-6-diamidino-2-phenylindole (DAPI) for 5 min and washed 3 times before mounting for confocal analysis using a Leica TCS SP2 laser-scanning confocal microscope.

Generation of rabbit anti-ZIP8 antibody—Anti-ZIP8 antiserum was generated in rabbit against peptide (CDHTHFRNDDFGSKEK) that is conserved between mouse and rat ZIP8. Anti-ZIP8 antibodies were affinity purified by using SulfoLink Coupling Resin (Thermo Scientific).

Iron loading and isolation of cell surface proteins—To load cells with iron, H4IIE cells were treated with 100 or 250 µM ferric nitrilotriacetic acid (NTA) for 72 h. Cell surface proteins were isolated by using the Pierce® Cell Surface Protein Isolation Kit (Thermo Scientific) according to the manufacturer’s protocol. Briefly, H4IIE cells were incubated with EZ-Link Sulfo-NHS-SS-Biotin to biotinylate cell surface proteins, which were subsequently affinity purified by using NeutrAvidin Agarose resin (Thermo Scientific). Bound proteins were released by incubating with SDS-PAGE buffer containing 50 mM DTT.

Site-directed mutagenesis of potential N-linked glycosylation sites in rat ZIP8—The QuickChange® Lightning Site-Directed Mutagenesis Kit (Stratagene) was used to mutate in rat ZIP8 asparagine (N) residues at positions 40, 88, and 96 to aspartic acid (D) or glutamine (Q). Residues were mutated individually or in combination. All constructs were verified by DNA sequencing.

Assessment of N-linked glycosylation—Cell lysates were digested with PNGase F to cleave N-linked glycans. Samples (30 µg protein) were denatured in buffer containing 1% (v/v) 2-mercaptoethanol and 0.5% (w/v) SDS at 37 °C for 30 min and then digested for 2 h at 37 °C with PNGase F (50,000 unit/mL of sample volume or 50 units/µg protein) (New England Biolabs). To inhibit N-glycosylation of endogenous ZIP8, H4IIE cells were incubated with tunicamycin (2 µg/mL) for 48 h.

Western blot analysis—Proteins were mixed with Laemmli buffer, incubated at 37 °C for 20 min, electrophoretically separated on a 7.5% polyacrylamide gel, and transferred to a nitrocellulose membrane (Schleicher and Schuell). The blot was blocked for 1 h in 5% (w/v) nonfat dry milk in Tris-buffered saline containing 0.1% (v/v) Tween 20 (TBST). Blots were then incubated overnight at 4 °C or 2 h at RT in blocking buffer containing 2.0 µg/ml mouse anti-HA (Roche) or affinity-purified rabbit anti-ZIP8, mouse anti-transferrin receptor 1 (TFR1) (1:4000; Invitrogen), rabbit anti-scavenger receptor B1 (SR-B1) (1:2000; Novus Biologicals), mouse anti-tubulin (1:10,000; Sigma), or rabbit anti-CCS (1:5000; Santa Cruz Biotechnology). The membrane was washed 4 times for 5 min with TBST and incubated with appropriate HRP-conjugated secondary antibodies. Immunoreactivity was visualized by using enhanced chemiluminescence (SuperSignal West Pico, Pierce) and X-ray film or the FluorChem E digital darkroom (Cell Biosciences).

Measurement of pH-dependent iron transport activity in HEK 293T cells—The pH-dependent iron transport activity was determined as previously described (20). Briefly, the uptake buffer (130 mM NaCl, 10 mM KCl, 1 mM CaCl2 and 1 mM MgSO4) was adjusted to pH 7.5, 6.5, and 5.5 by using HEPES (4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid) or MES (2-(N-morpholino)ethanesulfonic acid) buffers (2). HEK 293T cells transiently transfected for 48 h with mouse ZIP8 or empty vector (pCMV-Sport6) were washed twice with SFM and then incubated for 1 h in SFM containing 2% (w/v) BSA at 37 °C in 5% CO2. For uptake, cells were incubated with 2 µM 59Fe-ferric citrate in uptake buffer in the presence of 1 mM L-ascorbic acid at 37 °C for 1 h.

Measurement of mRNA levels—Total RNA from H4IIE cells was isolated by using RNA Bee (Tel-Test) according to the manufacturer’s protocol. Relative mRNA levels of ZIP8, TFR1, and cyclophilin B in H4IIE cells were determined as described previously (22). Transcript copy numbers of ZIP8, ZIP14, DMT1, and TFR1 mRNA in human RNA (Human Total RNA Master Panel II, Clontech)
were determined by using quantitative RT-PCR and standard curves generated from the plasmids pCMV-Sport6–human ZIP8 (BC012125; Open Biosystems), pCMV-XL4–human ZIP14 (BC015770; Open Biosystems); pBluescriptR–human DMT1 (BC100014, Open Biosystems); and pOTB7–human TFR1 (BC001188; ImaGenes). TaqMan primers and probes (Applied Biosystems) used for PCR were selected to target all known mRNA variants.

Suppression of ZIP8 expression in BeWo cells and measurement of iron uptake—BeWo cells were reverse transfected with 50 nM siRNA targeting ZIP8 mRNA (FlexiTube siRNA, Qiagen) or AllStars Negative Control siRNA (Qiagen) by using Lipofectamine RNAi Max (Invitrogen). After 72 h, cell lysates were prepared in RIPA buffer and analyzed by Western blotting for ZIP8 as described above. For primary antibody, rabbit anti-human SLC39A8, Prestige Antibodies (Sigma-Aldrich) was used at 1:5000 dilution. Iron uptake was performed as described for HEK293 T cells.

Statistical and regression analysis—Statistical and regression analyses were performed using SigmaPlot version 12 (Systat Software) with critical significance level $\alpha = 0.05$. Data for radiotracer uptake in oocytes were analyzed using two-way analysis of variance (ANOVA) followed by pairwise multiple comparisons using the Holm–Šidák test (adjusted $P$ is reported). Data were fit by Equation 1 by using the least-squares method of regression followed by $F$-tests of the significance of the fit to the model; SE is the standard error of the estimate, and $P$ is the significance of the fit.

RESULTS

Iron and zinc transport by ZIP8—Overexpression of ZIP8 in HEK 293T cells (Fig. 1A) was associated with a 200% increase in the uptake of $^{59}$Fe compared with that in control cells transfected with empty vector (Fig. 1B). Consistent with the reported role of ZIP8 as a zinc transporter, cells transfected with ZIP8 cDNA displayed a 40% increase in the uptake of $^{65}$Zn (Fig. 1B). The uptake of $^{59}$Fe by ZIP8-expressing cells was inhibited $>90\%$ by a 10-fold molar excess of zinc (Fig. 1C). A 10-fold excess of iron likewise inhibited $^{65}$Zn uptake. Uptake studies for iron were conducted in the presence of ascorbate so that uptake of Fe$^{2+}$ could be directly assessed. Without ascorbate, iron uptake from ferric citrate requires a reduction step (likely via a cell surface ferrireductase), which could affect uptake kinetics.

Metal-ion substrate profile of ZIP8 expressed in Xenopus oocytes—Expression of rat ZIP8 strongly stimulated the uptake of 5 µM $^{109}$Cd$^{2+}$, $^{57}$Co$^{2+}$, $^{55}$Fe$^{2+}$, and $^{65}$Zn$^{2+}$ and, more weakly, the uptake of $^{54}$Mn$^{2+}$ in Xenopus oocytes at pH 7.5 (Fig. 2A). We found also that expression of mouse ZIP8 in oocytes similarly stimulated the uptake of 2 µM $^{55}$Fe$^{2+}$ ($P < 0.001$, data not shown). Expression of rat ZIP8 did not stimulate the transport of $^{64}$Cu$^{2+}$ or $^{65}$Cu$^{+}$ (Fig. 2A). ZIP8-mediated $^{57}$Co$^{2+}$ transport (after subtraction of $^{55}$Co$^{2+}$ uptake in control oocytes) was saturable (supplemental Fig. S1) with half-maximal Co$^{2+}$ concentration ($K_{0.5}^{Co}$) of $\approx 1.0$ µM. We have previously demonstrated that mouse ZIP14, a close homolog of ZIP8, transports Cd$^{2+}$, Fe$^{2+}$, Mn$^{2+}$, and Zn$^{2+}$ but not Cu$^{+}$ or Cu$^{2+}$ (9), and found, in the present study, that mouse ZIP14 also transported Co$^{2+}$ (supplemental Fig. S1). The substrate profile of ZIP8 therefore closely matches that of ZIP14.

Properties of ZIP8-mediated iron transport in oocytes—ZIP8-mediated transport of $^{55}$Fe$^{2+}$ was strongly inhibited by 10-fold excess Co$^{2+}$ and Zn$^{2+}$ and more modestly by Mn$^{2+}$ (Fig. 2B), this last observation consistent with the much lower transport activity we had observed for $^{54}$Mn$^{2+}$ relative to Co$^{2+}$ and Zn$^{2+}$. ZIP8-mediated transport of 2 µM $^{55}$Fe$^{2+}$ was markedly pH dependent with optimal pH $\approx 7.5$ (Fig. 2C). ZIP8 did not mediate significant $^{55}$Fe$^{2+}$ transport at pH 6.5 or below (Fig. 2C). Therefore ZIP8 is active over roughly the same pH range (for Fe$^{2+}$ transport) as is ZIP14 (2). ZIP8-mediated $^{55}$Fe$^{2+}$ transport (after subtraction of $^{55}$Fe$^{2+}$ uptake in control oocytes) was saturable and ZIP8 exhibited high affinity for Fe$^{2+}$ (Fig. 2D, E). The concentration at which Fe$^{2+}$ transport was half-maximal ($K_{0.5}^{Fe}$) was $\approx 0.7$ µM and did not significantly differ from $K_{0.5}^{Co}$ ($P = 0.27$, by Student’s $t$ test).
**Subcellular localization of ZIP8**—For localization studies, we constructed an expression vector encoding rat ZIP8 with a C-terminal HA epitope tag. Western blot analysis of lysates from cells transfected with ZIP8 or ZIP8-HA cDNA revealed a similar band pattern when probed with anti-ZIP8 antibody (Fig. 3A). Two main immunoreactive bands were detected, one at ~100 kDa and the other at ~45 kDa. The latter band is similar to the 50.2 kDa theoretical molecular weight of rat ZIP8 (NCBI Reference Sequence NP_001011952.1), calculated from the amino acid sequence of the nascent peptide by using the ExPASy Compute pI/Mw tool (http://web.expasy.org/cgi-bin/compute_pi/pi_tool). After the blot was stripped and re-probed with anti-HA antibody, the HA-tagged protein was detected whereas the untagged protein was not (Fig. 3A). Detection of the same-sized bands for ZIP8-HA with either the anti-ZIP8 antibody or anti-HA antibody confirms that the anti-ZIP8 antibody detects expressed rat ZIP8 protein.

To determine if ZIP8 is expressed at the plasma membrane, HEK 293T cells were transfected with ZIP8-HA, fixed, and analyzed by immunofluorescence microscopy. Cells were analyzed without or with the detergent saponin to permeabilize the plasma membrane. In nonpermeabilized cells, strong plasma membrane staining was observed for ZIP8 (Fig. 3B), consistent with an exofacial C-terminal HA tag. Treatment with saponin permeabilized the cells, as indicated by the detection of the intracellular protein LAMP-1. In permeabilized cells, ZIP8 was detected not only at the plasma membrane, but also widely throughout the cytosol in an apparent vesicular distribution. Colocalization studies (Fig. 3C) revealed that intracellular ZIP8 partially colocalized with EEA1, a marker of early endosomes. Since localization studies using overexpressed proteins can lead to artifacts, future studies will need to examine the localization of endogenous ZIP8.

**pH-dependent iron transport activity of ZIP8**—The detection of ZIP8 in early endosomes of HEK 293T cells led us to consider a role for ZIP8 in iron transport from the acidic endosomal compartment. We therefore assessed the effect of pH on iron transport by ZIP8. Iron transport in HEK 293T cells was highest at pH 7.5, and then decreased with decreasing pH (Fig. 4). ZIP8-transfected cells displayed enhanced iron uptake at pH 6.5 but not at pH 5.5.

**N-glycosylation of ZIP8**—Rat ZIP8 contains three potential N-linked glycosylation sites at residues 40, 88, and 96. We used site-directed mutagenesis along with Western blot analysis to investigate the glycosylation status of each of these sites. Mutation of each site independently (N40D, N88Q, or N96Q) increased the electrophoretic mobility of the expressed protein mutants compared to wild-type ZIP8 (Fig. 5A). The faster mobility of the mutant proteins is consistent with the prediction of fewer glycans. Accordingly, when two of the three sites were mutated in combination, the proteins migrated even faster, whereas the triple mutant protein (N40D/N88Q/N96Q) migrated the fastest. The immunoreactive band at ~100 kDa is approximately twice the size of the band at ~45 kDa, and both bands shifted down in parallel in the mutant proteins. These observations suggest that the ~100-kDa protein represents a dimer of the ~45-kDa protein. Iron transport studies with the triple mutant (N40D/N88Q/N96Q) protein revealed that N-linked glycans are not required for iron or zinc transport by ZIP8 (supplemental Fig. S4).

To determine if endogenous rat ZIP8 is glycosylated, we used H4IIE cells, a rat hepatoma cell line. We chose to examine a liver-derived cell line because hepatocytes are able to take up NTBI as well as assimilate iron from transferrin (23,24). The most prominent immunoreactive band in H4IIE lysates detected by anti-ZIP8 antibody migrated with an apparent molecular mass of ~140 kDa (Fig. 5B, lanes 1 and 2 and supplemental Fig. S2), similar to what others have shown in Western blot analyses of ZIP8 (25). ZIP8 was present at the cell surface because it could be isolated by using cell-surface biotinylation (Fig. 5B, lanes 3 and 4). Treatment of isolated cell-surface proteins with the endoglycosidase PNGase F shifted the cell-surface ZIP8 band from 140 kDa to ~80 kDa, indicating that endogenous ZIP8 in rat H4IIE cells contains N-linked glycans.

Treatment of cells with the glycosylation inhibitor tunicamycin eliminated the immunoreactive ZIP8 band at 140 kDa and increased the intensity of the band at 100 kDa.
(Fig. 5C, lanes 1 and 2 vs. 5 and 6), suggesting that the 100 kDa band is a less glycosylated form of ZIP8. Additional treatment of lysates of tunicamycin-treated cells with PNGase F reduced the intensity of the 100-kDa band and increased the intensity of the 80-kDa band (Fig. 5C, lanes 5 and 6 vs. 7 and 8). Collectively, these observations suggest that the 100-kDa band is a partially glycosylated form of ZIP8 whereas the 80-kDa band is deglycosylated. The immunoreactive band detected at ~150 kDa in H4IIE cells appears to be nonspecific because it is unaffected by PNGase F (Fig. 5C, lanes 1 and 2 vs. 3 and 4) or tunicamycin (Fig. 5C, lanes 3 and 4 vs. 7 and 8).

Iron loading and ZIP8 expression—Treatment of H4IIE cells with either 100 or 250 μM iron nitrilotriacetic acid (FeNTA) for 72 h increased ZIP8 levels 2- and 4-fold respectively as determined by Western blot analysis and densitometry (Fig. 6A). We used the level of Tfr1, known to be decreased under high-iron conditions (26), as an indicator of cellular iron loading in response to FeNTA treatment. Cell-surface biotinylation experiments revealed an increase in ZIP8 and a decrease in TFR1 abundance at the plasma membrane in cells treated with iron (Fig. 6B). The plasma membrane protein SR-B1 was measured to demonstrate enrichment of plasma membrane proteins isolated by cell-surface biotinylation as well as equivalent lane loading. The cytosolic protein CCS (copper chaperone for superoxide dismutase) was detected in total cell lysate but not in isolated cell-surface proteins, indicating that the isolated cell-surface extract was not contaminated with cytosolic proteins. The higher ZIP8 levels in iron-loaded cells appear to result from post-transcriptional events because ZIP8 mRNA levels did not increase in response to iron loading (Fig. 6C), whereas TFR1 mRNA levels, used as a positive control for iron-related changes in gene expression, were substantially decreased in iron-loaded cells. Treatment of cells with zinc (40 μM zinc sulfate) increased ZIP8 levels, but the effect was transient, occurring at 3 h after treatment and not after 48 h (supplemental Fig. S3). The higher ZIP8 levels at 3 h were not associated with higher ZIP8 mRNA levels (data not shown), suggesting that zinc, like iron, regulates ZIP8 in a post-transcriptional manner.

Effect of FeNTA treatment on cellular zinc concentrations and effect of zinc depletion on ZIP8 levels—Although treatment of cells with FeNTA increases ZIP8 levels, it is possible that this effect is mediated through a secondary zinc deficiency induced by the high-iron treatment. To examine this possibility, we treated cells with FeNTA and then measured total cellular iron and zinc concentrations by ICP-MS. As a positive control for zinc depletion, we included cells that were treated with the zinc-specific chelator, TPEN, which decreased zinc concentrations in H4IIE cells by 60%, but did not affect iron concentrations (supplemental Fig. 5A). We found that treatment of cells with 250 μM FeNTA increased cellular iron levels 12-fold and decreased zinc concentrations by 30% (supplemental Fig. 5A). To determine if zinc deficiency could independently increase ZIP8 levels, we measured ZIP8 levels in lysates from cells treated with TPEN. Western blot analysis revealed that zinc deficiency alone did not increase ZIP8 levels, whereas treatment with Fe did (supplemental Fig. 5B). We therefore conclude that ZIP8 levels increase in iron-treated cells in response to iron loading but not zinc depletion.

Tissue expression of ZIP8—To determine where ZIP8 is expressed, we measured mRNA copy number in RNA isolated from 20 different human tissues. Most abundant ZIP8 expression was detected in lung and placenta followed by salivary gland and thymus (Fig. 7). Tissues expressing the lowest amount of ZIP8 included skeletal muscle, fetal brain, testis, fetal liver, and small intestine. To compare the tissue distribution and expression levels of ZIP8 to other known proteins involved in cellular iron import, we measured mRNA copy numbers of ZIP14, DMT1, and TFR1. Similar to previous studies, ZIP14 mRNA levels were highest in liver and heart (4), DMT1 mRNA was abundantly expressed in kidney, brain, and thymus (2) and TFR1 was abundantly expressed in placenta (27).

Iron transport activity of endogenous ZIP8—To examine iron-transport activity of endogenous ZIP8, we measured iron uptake in BeWo cells in which ZIP8 was suppressed by
using siRNA. BeWo cells, a placental cell line derived from a human choriocarcinoma, are widely used as a model to study placental uptake of nutrients including iron (28). In BeWo cells treated with ZIP8 siRNA, ZIP8 protein levels were reduced by 90% compared to control cells treated with negative control siRNA (Fig. 8A). After suppression of ZIP8, iron uptake was reduced by 37% (Fig. 8B), suggesting that ZIP8 mediates, at least in part, iron uptake by BeWo cells.

DISCUSSION

The present study establishes that ZIP8 can mediate the cellular uptake of iron, making it the third mammalian transmembrane iron-import protein to be identified. ZIP8 was first identified in 2002 in a screen of monocyte cDNAs induced by infection and inflammatory stimuli (18). The amino acid sequence of ZIP8 was found to share significant homology to the zinc transporters ZIP1 and ZIP2, and therefore ZIP8 was postulated to function as a zinc transporter. Support for this possibility was provided by the observation that CHO cells stably transfected with ZIP8 accumulated and retained more zinc, as demonstrated by using an intracellular zinc-selective fluorescent dye (18). The first characterized mammalian iron importer, DMT1, was identified by screening an iron-deficient rat duodenal cDNA library for iron-transport activity in Xenopus oocytes (2). The second protein found capable of transporting iron into mammalian cells was ZIP14, which had been originally described as a zinc transporter (29). Iron uptake via ZIP14 was demonstrated in HEK 293T and Sf9 insect cells that overexpressed the protein and then later in the Xenopus oocyte expression system (5,9). Here we document the iron transport activity of ZIP8 by using both transfection of HEK 293T cells and expression in Xenopus oocytes.

Like ZIP14, we found that ZIP8 transports iron at physiologic pH, is expressed at the cell surface, and mediates the uptake of iron from ferric citrate, the predominant form of NTBI in the plasma of individuals with iron overload (30). Although these observations suggested a role for ZIP8 in NTBI uptake, we found that ZIP8 is not abundantly expressed (at least according to mRNA copy number) in liver, the primary organ that clears NTBI (31), or the heart, which also takes up NTBI (32). On the other hand, ZIP8 has been reported to be abundantly expressed in human pancreas (18), which was not represented in our RNA panel. After the liver, the pancreas is the organ that takes up the most NTBI (31). Our data in H4IIE hepatoma cells showing that iron loading increases plasma membrane levels of ZIP8 suggest that ZIP8 may be recruited to the cell surface to enhance the uptake of NTBI. Indeed a number of studies have shown that iron loading increases plasma membrane levels of ZIP8 (12). Iron-mediated cardiotoxicity is the main cause of death in β-thalassemia major patients, who eventually succumb to iron overload due to repeated blood transfusions (37). Interestingly, microarray analysis found that ZIP8 mRNA levels were 1.8-fold higher in cardiomyocytes from β-thalassemic mice compared with wild-type controls (38). By contrast, DMT1 mRNA levels in cardiomyocytes from β-thalassemic mice were downregulated 1.7 fold, whereas the levels of ZIP14 were apparently unaffected. Future studies should explore the contribution of ZIP8 to NTBI uptake in hepatocytes and cardiomyocytes as well as other cell types susceptible to iron overload.

Treatment of H4IIE cells with zinc increased ZIP8 levels (though transiently), suggesting that ZIP8 actively participates in cellular zinc acquisition. As a zinc transporter, ZIP8 is frequently cited as a lysosomal transporter (39,40), yet in most studies it was detected at the plasma membrane (reviewed in (16)). In a study using Madin-Darby canine kidney (MDCK) cells, ZIP8 abundance at the plasma membrane was found to be responsive to zinc, increasing with zinc depletion and returning to basal levels after zinc repletion (19). Therefore, although we found that zinc increases ZIP8 levels in H4IIE cells, it will be important to determine if the increase occurs at the plasma membrane. The upregulation of the zinc/iron importer ZIP8 in response to iron loading may partially account for hepatic zinc accumulation that occurs during iron overload (22,41-43).
We have previously observed for mouse ZIP14 a pattern of incomplete mutual inhibition and proposed a model in which transport activities of Cd\(^{2+}\) and Zn\(^{2+}\) differed from the transport activities of other metal ions (9). Whereas 10-fold excess Zn\(^{2+}\) entirely abolished \(^{55}\)Fe\(^{2+}\) uptake, Fe\(^{3+}\) only weakly inhibited ZIP14-mediated Zn\(^{2+}\) transport (9). In contrast, we found for ZIP8 that 10-fold excess Zn\(^{2+}\) did not abolish \(^{55}\)Fe\(^{2+}\) uptake in oocytes but inhibited \(^{55}\)Fe\(^{2+}\) uptake by 77% (Fig. 2B), i.e. to roughly the extent expected for a 10-fold excess of inhibitor, and we observed mutual inhibition between Fe\(^{2+}\) and Zn\(^{2+}\) in HEK 293T cells expressing ZIP8 (Fig. 1C). Therefore, our observations for ZIP8—but not ZIP14—are consistent with a single homogeneous transport pathway (3) shared by Fe\(^{2+}\) and Zn\(^{2+}\).

The iron-transport activity of ZIP8 along with its partial colocalization with EEA1 raises the possibility that ZIP8 may participate in the cellular assimilation of iron from the transferrin cycle. Specifically, ZIP8 could serve to transport iron from the acidified endosome into the cytosol, a function usually attributed to DMT1 (3), and more recently to ZIP14 (10). However, it should be noted that the iron transport activity of ZIP8 decreases with decreasing pH. At pH 6.5, the pH by which ~50% of the iron dissociates from transferrin-decreasing pH. At pH 6.5, the pH by which ~50% of the iron dissociates from transferrin receptor complex in endosomes (44), ZIP8 showed some iron transport activity in HEK 293T cells but not in oocytes. At pH 5.5, often found in lysosomes, ZIP8 showed no iron transport activity. Thus, it seems unlikely that ZIP8 would function as an iron transporter in lysosomes, where it has been detected (18,45). Lysosomes play a key role in cellular iron metabolism by degrading the iron-storage protein ferritin and releasing the iron into the cytosol, especially during situations such as iron deprivation (46). How iron is transported across the lysosomal membrane remains unknown.

Although the substrate profile of ZIP8 and ZIP14 are similar, they exhibit markedly different tissue expression profiles, suggesting nonredundant functions. In agreement with other studies, we found most abundant ZIP8 expression in the lung (18,47,48). The lung is not a significant portal of entry for iron, but other metals such as cadmium and manganese can enter the body in toxic amounts through the lungs (49). It seems clear from our data and others (19) that ZIP8 expressed in oocytes transports cadmium, but in contrast to a previous study (19), we found that ZIP8 is not an efficient manganese transporter. Indeed our data overall do not support a previous conclusion that ZIP8 acts principally as a manganese transporter under physiologic conditions (19) since (i) manganese transport activity of ZIP8 was very low relative to other metal ions that are more abundant in serum (namely iron and zinc); and (ii) we obtained \(K_{0.5}\) for iron of 0.7 \(\mu\)M, substantially lower than that reported for manganese (2.2 \(\mu\)M) (19), suggesting that ZIP8 has a higher affinity for iron than for manganese.

After the lung, the tissue we found that expressed the most ZIP8 was placenta. Strong ZIP8 expression in placenta was also noted in a previous study in which placental ZIP8 expression was second highest out of 16 human tissues after pancreas (and even more abundant than in lung) (18). It is unknown how iron is transported across the placenta to the developing fetus (50). We do know, however, that DMT1 and ZIP14 are not required for maternofetal iron transfer because Dmt1 and Zip14 (Slc39a14) knockout mice are born alive with adequate amounts of iron (50). These observations leave ZIP8 as the only remaining known iron transport protein that could function in the placenta, although additional yet-to-be identified iron transporters may exist.

While our manuscript was under review, Galvez-Peralta et al. (52) reported the characterization of the ZIP8 hypomorph mouse, which carries a neomycin-resistance gene in intron 3 of Zip8 (Slc39a14), resulting in markedly decreased levels of ZIP8 in the embryo, yolk sac, fetus, placenta, and other tissues of neonates. ZIP8 hypomorph mice exhibit severe anemia as embryos and neonates and do not survive more than 48 hours after birth. The anemia appears to result from iron deficiency as indicated by low serum iron, total iron-binding capacity, and low hepatic iron. The iron-deficient phenotype of the hypomorph mice strongly suggests that ZIP8 plays a direct physiologic role in iron metabolism. At the cellular level, ZIP8 seems directly involved, as iron uptake from fetal...
fibroblasts and liver cells from the hypomorphic mice was found to be significantly impaired. In the ZIP8 hypomorphs, the lower levels of hepatic iron suggests that placental iron transfer is impaired. Consistent with this possibility is our observation that siRNA-mediated knockdown of endogenous ZIP8 reduces iron uptake by BeWo cells, a widely used in vitro model system for placental iron transport (28).

The demonstration that ZIP8—and previously ZIP14 (5)—can transport iron raises the question as to whether iron transport activity is a general characteristic of the ZIP family of proteins. In initial screens of the remaining 12 ZIP family members, we have evidence that only one of these, ZIP2, appears capable of transporting iron (unpublished observations W. Zhang and M. Knutson). Interestingly, previous studies of the Zip2 knockout mouse concluded that ZIP2 plays a role in iron homeostasis (53). In an effort to identify which of the 14 ZIP family members might transport iron, we previously examined the effect of iron deficiency and iron overload on the expression of all the ZIP proteins in rat liver (22). We reasoned that iron-regulated ZIP proteins represented the best candidates for possible iron transport activity. However, in that study, ZIP8 mRNA levels did not vary with iron status in rat liver; nor did they vary in H4IIE cells treated with FeNTA. Here we also find that ZIP8 mRNA levels did not vary with iron loading but ZIP8 protein levels did increase, suggestive of post-transcriptional regulation. These observations highlight the limitation of screening ZIP transporters at the mRNA level.

ZIP proteins generally are predicted to have a long extracellular N-terminal domain, sometimes comprising more than half of the amino acid sequence (54). The N-terminal region, which contains several conserved features including multiple predicted N-linked glycosylation sites, has been proposed to modulate metal transport activity (54). The presence of N-linked glycosylation has been demonstrated for mouse ZIP4 (55), ZIP5 (56), ZIP8 (19), ZIP14 (57) and human ZIP14 (4) by using the endoglycosidase PNGase F or the glycosylation inhibitor tunicamycin. Here we used PNGase F and tunicamycin to show that rat ZIP8 is glycosylated. Moreover, by using site-directed mutagenesis, we demonstrate that all three predicted N-linked glycosylation sites of rat ZIP8 are modified by glycans. Importantly, we found that these N-linked glycans are not required for iron or zinc transport by ZIP8 (supplemental Fig. S4). To our knowledge, this is the first direct assessment of the role of glycosylation on metal transport by a ZIP protein.

By using the Xenopus oocyte expression system, we found that ZIP8—and ZIP14—efficiently transport cobalt. Indeed, the substrate profiles of ZIP8 and ZIP14 are similar to the ZIP protein ZUPT, which transports zinc, iron, cobalt, and possibly manganese and cadmium in E. coli (14). Little is known about mammalian cobalt transport, although some evidence indicates that DMT1 (58,59) and its homologue Nramp1 (58) can transport cobalt. Intestinal absorption of cobalt has been shown to increase in iron-deficient animals and humans (60,61), an effect likely mediated through DMT1, which is markedly upregulated in iron-deficient intestine (2). By contrast, it seems unlikely that DMT1 mediates the uptake of cobalt by the liver because hepatic cobalt concentrations are unaffected in mice with hepatocyte-specific deletion of Dmt1 (unpublished observations, C. Wang and M. Knutson). Our tissue expression profile data implicate ZIP14 as a likely candidate for hepatic cobalt uptake since liver ZIP14 mRNA levels are approximately 10 times higher than DMT1 or ZIP8. In the lung, ZIP8 expression is markedly higher than ZIP14 or DMT1, and therefore ZIP8 may represent a major cobalt transporter in this tissue. It is well known that cobalt is absorbed via the lungs (62), which can result in cobalt toxicity, such as in workers occupationally exposed to cobalt-containing dusts. It should be noted, however, that under normal circumstances, the main source of cobalt exposure is food; yet the amount (as inorganic, non-cobalamin cobalt) in typical human diets (< 40 μg/d) is markedly lower than the usual amount of iron, zinc, or manganese (2-20 mg/d). In serum, cobalt concentrations (~ 0.2 ng/mL) are typically less than manganese (~ 0.6 ng/mL) and at least 3 orders of magnitude lower than those of iron or zinc (~ 1 μg/mL) (63). Therefore, the principal physiologic substrates of ZIP8 and ZIP14 are
likely to be iron and zinc, and to a lesser extent, manganese and cobalt.

In conclusion, ZIP8 is a novel iron transporter whose expression profile suggests that it may play a role in iron metabolism in the placenta, pancreas, lung, and other organs. The upregulation of cell-surface ZIP8 by iron loading additionally suggests that ZIP8 may contribute to NTBI uptake during iron overload. Future studies of Zip8 knockout animals, especially conditional knockouts, will be required to define the role(s) of ZIP8 in iron metabolism in vivo.

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FOOTNOTES

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1To whom correspondence may be addressed: Food Science and Human Nutrition Department, PO Box 110370, University of Florida, Gainesville, FL 32611; Fax: 352-392-9467; E-mail: mknutson@ufl.edu

2Department of Molecular & Cellular Physiology, University of Cincinnati College of Medicine, Cincinnati, Ohio, 45267.

3Systems Biology & Physiology Program, University of Cincinnati College of Medicine, Cincinnati, Ohio, 45267.

4The abbreviations used are: ZIP, ZRT/IRT-like Protein; DMT1, divalent metal-ion transporter-1; NTBI, non-transferrin-bound iron; EEA1, early endosome antigen 1; LAMP-1, lysosome associated macrophage protein-1; TFR1, transferrin receptor 1.

FIGURE LEGENDS

FIGURE 1. Overexpression of ZIP8 increases the cellular uptake of iron and zinc. A, Western blot analysis of ZIP8 in HEK 293T cells transiently transfected with empty pExpress vector or pExpress-rat ZIP8. Total cell lysates were resolved by SDS-PAGE, transferred to nitrocellulose membrane, and immunoblotted with affinity-purified anti-ZIP8 antibody. B, Cellular uptake of iron and zinc in HEK 293T cells overexpressing ZIP8. Forty-eight hours after transfection, cells were incubated for 1 h in uptake medium containing 2 µM $^{59}$Fe-ferric-citrate or $^{65}$Zn-ZnCl$_2$, and cellular uptake of $^{59}$Fe (left) or $^{65}$Zn (right) was measured by gamma counting. C, Mutual inhibition of iron and zinc uptake in cells overexpressing ZIP8. $^{59}$Fe and $^{65}$Zn uptake were measured in the presence of a 10-fold molar excess of unlabeled zinc (left) or iron (right). The amount of $^{59}$Fe or $^{65}$Zn taken up by cells is expressed as pmol/mg protein. Data represent the mean ± S.E. of three independent experiments. Treatment group means were compared by unpaired Student’s t test.
FIGURE 2. Functional properties of ZIP8 expressed in Xenopus oocytes. A, Metal-ion substrate profile of rat ZIP8. Transport (uptake) of 5 µM metal ions (*M") was measured at pH 7.5 in control oocytes and oocytes expressing rat ZIP8. Data are mean and SD, n = 14–19 oocytes per group. Two-way ANOVA revealed an interaction (P < 0.001); Holm–Šidák pairwise comparisons within each metal revealed that uptake of Cd²⁺, Co²⁺, Fe²⁺, Mn²⁺ and Zn²⁺ differed between ZIP8 and control (P < 0.001 except for Mn²⁺, P = 0.016), whereas uptake of Cu¹⁺ (P = 0.81) and Cu²⁺ (P = 0.47) did not differ between ZIP8 and control. B, Inhibition profile of ZIP8-mediated ⁵⁵Fe²⁺ transport. Transport of 1 µM ⁵⁵Fe²⁺ in control oocytes (gray bars) and in oocytes expressing ZIP8 (black bars) at pH 7.5 in the absence of inhibitor ('None') or in the presence of 10 µM Co²⁺, Mn²⁺ or Zn²⁺. Data are mean and SD, n = 11–13 oocytes per group. Two-way ANOVA revealed an interaction (P < 0.001); within ZIP8, all inhibitor metal ions differed from 'None' (P < 0.001). C, Transport of 2 µM ⁵⁵Fe²⁺ as a function of extracellular pH in control oocytes (gray circles) and oocytes expressing ZIP8 (black circles). Data are mean and SD, n = 8–13 oocytes per group. Two-way ANOVA revealed an interaction (P < 0.001); ZIP8 differed from Control at pH 7.0 and higher (P < 0.001) but not at pH 6.5 and lower (P ≥ 0.22); within ZIP8, transport differed for every pairwise comparison (P < 0.008) except between pH 7.0 and 8.0 (P = 0.94) and between pH 6.0 and 5.5 (P = 0.87). D,E, Saturation kinetics of ZIP8-mediated ⁵⁵Fe²⁺ transport. Uptake of ⁵⁵Fe²⁺ was measured in control oocytes (gray circles) and oocytes expressing ZIP8 (black circles) at pH 7.5 (in D); data are mean and SD, n = 10–15 oocytes per group. Mean uptake in control oocytes was subtracted from mean uptake in oocytes expressing ZIP8 to obtain mean ZIP8-mediated ⁵⁵Fe²⁺ transport (in E) ± propagated standard error. ZIP8-mediated ⁵⁵Fe²⁺ transport data were fit by Equation 1 (solid black line) yielding \( V_{\text{Fe}}^\text{max} = 0.065 \pm 0.004 \text{ pmol.min}^{-1} \) and \( K_{\text{Fe}}^\text{0.5} = 0.66 \pm 0.16 \mu\text{M} \) (adjusted \( r^2 = 0.94; P < 0.001 \)).

FIGURE 3. ZIP8 expressed in HEK 293T cells localizes to the plasma membrane and intracellular vesicles. A, Western blot analysis of ZIP8 and ZIP8-HA. HEK293T cells were transiently transfected with empty pExpress vector, pExpress-rat ZIP8, or pExpress-rat ZIP8-HA. Total cell lysates were resolved by SDS-PAGE, transferred to nitrocellulose membrane, and immunoblotted with affinity-purified anti-ZIP8 antibody (left). The blot was then stripped and reprobed with an anti-tubulin antibody to indicate lane loading (right). B, Detection of ZIP8 in non-permeabilized and permeabilized cells. HEK 293T cells transiently transfected with pExpress empty vector or vector encoding ZIP8-HA were fixed and incubated without (-) or with (+) saponin to permeabilize cells. To detect ZIP8-HA, cells were incubated with anti-HA primary antibody followed by Alexa Fluor 594-conjugated secondary antibody (red). As a control for permeabilization, all cells were co-incubated with anti-LAMP1 primary antibody followed by Alexa Fluor 488-conjugated secondary antibody (green). Cells were analyzed by confocal fluorescence microscopy. C, Colocalization of ZIP8 and EEA1. HEK 293T cells transiently transfected with pExpress empty vector or vector encoding ZIP8-HA were fixed, incubated with saponin, and analyzed for ZIP8 as described in B. To detect EEA1, cells were co-incubated with anti-EEA1 primary antibody followed by Alexa Fluor 488-conjugated secondary antibody (green). Merged images of ZIP8 and EEA1 reveal areas of colocalization (yellow), indicated by arrows. Blue color indicates nuclei counterstained with DAPI.

FIGURE 4. pH dependence of ZIP8-mediated iron transport. HEK 293T cells were transfected with empty pCMV-Sport6 vector or pCMV-Sport6-mouse ZIP8. Forty-eight h after transfection, cells were incubated with 2 µM ⁵⁵Fe-ferric citrate for 1 h in uptake buffer at pH 5.5, 6.5 and 7.5. The amount of ⁵⁵Fe taken up by cells is expressed as pmol/mg of protein. Data represent the mean ± S.E. of three independent experiments.

FIGURE 5. Mutational analysis of potential N-linked glycosylation sites of rat ZIP8 expressed in HEK cells and glycosylation analysis of endogenous ZIP8 in H4IIE rat hepatoma cells. A, Western blot analysis of wild-type (WT) rat ZIP8 and rat ZIP8 asparagine mutants expressed in HEK 293T cells. Asparagine (N) residues 40, 88, and 96 of rat ZIP8 were mutated individually or in combination to
aspartic acid (D) or glutamine (Q).  

B, Western blot analysis of cell-surface ZIP8 in H4IIE cell lysates treated without (-) or with (+) PNGase F. Cell-surface proteins were labeled with Sulfo-NHS-SS-Biotin and affinity purified by using streptavidin agarose prior to Western analysis. C, Western blot analysis of endogenous ZIP8 in H4IIE cells/cell lysates treated without (-) or with (+) tunicamycin or PNGase F. In panels B and C, lane numbers are indicated below the blots.

FIGURE 6. Iron loading increases ZIP8 levels in H4IIE rat hepatoma cells. A, Western blot analysis of ZIP8 in cells treated for 72 h with 0, 100, or 250 µM FeNTA. Total cell lysates were resolved by SDS-PAGE, transferred to nitrocellulose membrane, and immunoblotted with anti-ZIP8 antibody. The blot was then stripped and reprobed with an anti-TFR1 antibody followed by an anti-tubulin antibody to indicate lane loading. Relative protein levels of ZIP8 and TFR1, normalized to levels of tubulin, were determined by densitometry. B, Western blot analysis of ZIP8, TFR1, SR-B1, and CCS in total cell lysate and cell-surface proteins isolated from cells treated for 72 h with 0, 100, or 250 µM FeNTA. Cell-surface proteins were labeled with Sulfo-NHS-SS-Biotin and affinity purified by using streptavidin agarose prior to Western analysis. Relative cell-surface protein levels of ZIP8 and TFR1, normalized to levels of SR-B1, were determined by densitometry. C, Relative mRNA levels of ZIP8 and TFR1 were determined by using quantitative RT-PCR. Data represent the mean ± S.E. of three independent experiments. Treatment group means were compared by one-way ANOVA.

FIGURE 7. Tissue expression of ZIP8, ZIP14, DMT1 and TFR1. Tissue expression profiles were determined by using the Human Total RNA Master Panel II derived from 20 different tissues (Clontech). Transcript copy numbers were determined by using quantitative RT-PCR.

FIGURE 8. Suppression of ZIP8 expression decreases iron uptake by BeWo cells. A, Western blot analysis of lysates (50 µg protein) from BeWo cells treated with negative control (NC) siRNA or siRNA targeting ZIP8 mRNA. As a positive control for ZIP8, lysates (5 µg protein) were also analyzed from BeWo cells transfected with pCMV-Sport6-human ZIP8 cDNA (ZIP8) or empty pCMVSport6 vector (Vector) B, Cellular ⁵⁹Fe uptake in BeWo cells transfected with negative control (NC) siRNA or siRNA targeting ZIP8 mRNA. Data represent the mean ± S.E. of three independent experiments.
Figure 1

A

![Western blot image with Anti-ZIP8 antibody showing bands at Vector and ZIP8 levels](image)

B

![Bar graphs showing 59Fe uptake (pmol/mg protein) and 65Zn uptake (pmol/mg protein) for Vector and ZIP8 conditions with statistical significance](image)

C

![Bar graphs showing 59Fe uptake (pmol/mg protein) and 65Zn uptake (pmol/mg protein) for -Zn and +Zn conditions with statistical significance](image)
Figure 2
Figure 3
Figure 4

The diagram shows the uptake of $^{59}$Fe (pmol/mg protein) at different pH levels: 5.5, 6.5, and 7.5, comparing Vector and ZIP8 treatments. The data is represented with error bars indicating variability. Statistical significance ($P < 0.05$) is indicated between the treatments at pH 6.5 and 7.5, with ZIP8 showing higher uptake compared to Vector.
Figure 5

A

B

PNGase F
- - - - + +
Biotinylation
- - + + + +

C

Tunicamycin
- - - - + + + +
PNGase F
- - + + - - + +

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Figure 6

A

| FeNTA (μM) | 0  | 100 | 250 |
|------------|----|-----|-----|
| ZIP8       |    |     |     |
| TFR1       |    |     |     |
| Tubulin    |    |     |     |

B

| FeNTA (μM) | 0  | 100 | 250 |
|------------|----|-----|-----|
| Total Cell Lysate |    |     |     |
| Cell Surface |    |     |     |
| ZIP8       |    |     |     |
| TFR1       |    |     |     |
| SR-B1      |    |     |     |
| CCS        |    |     |     |

C

| FeNTA, μM | 0  | 100 | 250 |
|-----------|----|-----|-----|
| ZIP8      |    |     |     |
| TFR1      |    |     |     |

Relative protein levels

- ZIP8
- TFR1

Relative cell-surface protein levels

- ZIP8
- TFR1

P < 0.001
P < 0.05
Figure 7
ZIP8 Is an Iron and Zinc Transporter Whose Cell-surface Expression Is Upregulated by Cellular Iron Loading
Chia-Yu Wang, Supak Jenkitkasemwong, Stephanie Duarte, Brian Sparkman, Ali Shawkii, Bryan Mackenzie and Mitchell D. Knutson

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