Identification of a mammalian silicon transporter

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Silicon (Si) is the second most abundant element in the Earth’s crust and is ubiquitous in the diet, but the role it plays in mammalian physiology remains unclear. There is substantial evidence for its importance in the normal health and development of bone and connective tissues of vertebrates (6, 25, 43, 45), but a specific physiological and/or metabolic function has not been identified. In particular, proteins responsible for Si transport in mammals remain elusive. Silicon is essential for many algae (e.g., diatoms) to produce their exoskeleton and to complete their cell cycle (5, 21). It is also important in many species of plants, with both structural and physiological roles identified (12, 13, 27).

The first Si transporter to be identified (C/SIT1) was in the diatom species *Cylindrotheca fusiformis* (22), and SITs are now known from a wide range of diatoms (51), choanoflagellates (32), and haptophytes (11). In plants, Si transport occurs through a collaboration of two individual transporter types, one of which is responsible for influx (movement of Si into cell) and the other for efflux (movement of Si out of cell). Influx occurs through an aquaporin channel (e.g., Lsi1, Lsi6) whereas efflux occurs through an energy-dependent active transport process driven by a proton gradient (e.g., Lsi2) (29, 30). Despite the characterization of multiple Si transporters in algae and plants as described, no Si-transporting homologs have been reported in mammals yet (29, 30, 32).

Previously, we reported that Sprague-Dawley rats on a Si-depleted diet massively reduced their urinary Si output to maintain serum and tissue Si levels (24). This was at the expense of phosphorus, which was decreased in serum and bone (24). These findings suggested that the kidney may be actively involved in Si conservation under chronic Si deprivation and that, somehow, phosphate is lost in the process. Here we report on the mammalian phosphate transport protein Scl34a2, which was upregulated in the kidney of the rats deprived of dietary Si. This protein was found to induce Si efflux activity when expressed in *Xenopus* oocytes and to exhibit structural similarity with Lsi2 in many plants. Identification that Scl34a2 can transport Si provides new evidence...
for a biological role for this element in mammals and establishes another distinct gene family of Si transporters.

**MATERIALS AND METHODS**

**Silicon Depletion Study**

Kidneys were obtained from the study of Judahsingesh et al. (24) following 6 mo of dietary Si intervention. Three-week-old female Sprague-Dawley rats were maintained for 26 wk on a formulated low-Si feed (~3 µg Si/g feed), with either low-Si drinking water (~15 mg Si/g water; Si deplete group, n = 20) or with orthosilicic acid (H3SiO4) supplemented in the drinking water (53 µg Si/g water; Si replete group, n = 10). A reference group of rats received a normal laboratory maintenance chow diet (B&K Rat and Mouse Standard Diet; B&K Universal) which is naturally high in Si (322 µg Si/g feed) and tap water (5 µg Si/g water); see reference 24 for diet compositions. This third group of rats is referred to as the Si-high reference group. Total Si intakes were 0.17 mg Si kg body wt and tap water (5 µg Si/g water), with either low-Si drinking water (~15 mg Si/g water; Si deplete group, n = 20) or with orthosilicic acid (H3SiO4) supplemented in the drinking water (53 µg Si/g water; Si replete group, n = 10). A reference group of rats received a normal laboratory maintenance chow diet (B&K Rat and Mouse Standard Diet; B&K Universal) which is naturally high in Si (322 µg Si/g feed) and tap water (5 µg Si/g water); see reference 24 for diet compositions. This third group of rats is referred to as the Si-high reference group. Total Si intakes were 0.17 mg Si kg body wt -1 day -1 in the Si deplete group, 4.1 mg Si kg -1 day -1 in the Si replete group, and 18.5 mg Si kg -1 day -1 in the Si-high reference group. After 26 wk, rats were euthanized by asphyxiation with carbon dioxide gas as previously described (24). Rats were killed and processed one at a time, with one rat from each group on the same day. Tissues were then harvested, as previously described (24), and stored at -20°C immediately following harvesting and then at -80°C for long-term storage. All groups of rats and their tissues were treated in precisely the same fashion. Spot urine samples were collected from fasted rats (24). Urinary Si and P analysis was by inductively coupled plasma optical emission spectrometry (ICP-OES), as described below, and data were corrected for urinary creatinine (24). As previously described (24), all animal procedures were carried out in accordance with the UK Home Office Animal Scientific Procedures Act 1986 (Scientific Procedures on Living Animals). Use of laboratory animals was approved by King's College London (UK) Animal Ethics Committee and the UK Home Office. For this study, the left kidneys from n = 10 Si deplete, n = 8 Si replete, and n = 5 Si-high reference rats were ground in liquid nitrogen and total RNA was extracted with the Qiagen RNeasy Maxi kit for microarray and quantitative PCR analysis. This part of the study was carried out in 2008.

**Gene Array Analysis**

Five micrograms total RNA per sample were hybridized to Affymetrix GeneChip Rat Genome 230 2.0 arrays (n = 4 Si deplete and n = 4 Si replete kidneys). Gene chip robust multiarray analysis (gerMA) was used to normalize the data including a summarization step based on m-estimator values for the probe sets (58). Modified T-statistics were used to calculate significance of differential gene expression (10, 44) between the Si replete vs. Si deplete groups. Genes were selected as “differentially expressed” when false discovery rate q was < 0.1 (49). [The microarray data set has been submitted to the NCBI Gene Expression Omnibus: accession no: GSE58404.] }

**Expression Studies**

Quantitative real-time PCR was used to investigate the expression of relevant transcripts, including that of an internal control (18S), in the full cohort of rat kidney RNAs (n = 10 Si deplete, n = 8 Si replete, and n = 5 Si-high reference group). Transcripts were amplified with the TaqMan Universal protocol for real-time RT-PCR. The primers were TaqMan Gene Expression Assays consisting of a FAM reporter and TaqMan MGB probes. Differences in gene expression between groups were statistically analyzed by unpaired t-test.

**Immunohistochemistry**

Kidneys from a normal laboratory maintenance chow fed rat were excised immediately after necropsy and then fixed in 4% PBS-buffered paraformaldehyde. The samples were then cryo-protected via sucrose gradient and snap-frozen in iso-pentane cooled on dry ice. The frozen samples were then embedded in Optimal Cutting Temperature compound (VWR). Tissue sections were subsequently cryo-sectioned at 12 µm thickness, collected on SuperFrost slides (Thermo Scientific), and allowed to air dry overnight at room temperature. Sections were blocked with normal serum in PBS. Samples were then incubated with primary antibody against Slc34a2 (Genetex) or an appropriately matched isotype control. Primary antibody was then detected by incubation with goat anti-rabbit IgG (H+L) Alexa Fluor 488 conjugate (Invitrogen) before counterstaining the nuclei with Hoechst 33342 (Invitrogen) and the cytoskeleton (f-actin) with phallolidin CF633 (Biotium). Imaging was carried out on a Leica SP2 confocal microscope using a 1.2 numerical aperture ×63 water immersion lens. Images were collected using the Leica Application Suite software. Alongside Slc34a2 antibody-stained sections, images of the isotype controls were also collected under identical settings and in “matched” parallel tissue sections. A threshold removing any minor nonspecific signal in the isotype controls was then defined, with this threshold subsequently applied identically across all collected images to robustly identify Slc34a2. Staining for Slc34a2 was distinctly punctate so, as well as presentation in as-collected intensity format, images are also presented in binary format (i.e., all Slc34a2 signal that is brighter than isotype threshold shown at maximum intensity). This “view” was included to facilitate visualization of Slc34a2 locality within the limits of the printed image size.

**Urinary P and Si Analyses**

Fasting spot urine samples collected from 6-h fasted rats (n = 8 Si deplete, n = 5 Si replete, and n = 6 Si-high reference rats) were digested (in 1:1 mixture of 69% nitric acid and 40% hydrogen peroxide), diluted (1:100), and analyzed for total phosphorus by inductively coupled plasma optical emission spectrometry (ICP-OES; Jobin Yvon 2000-2) at 214.914 nm with sample-based standards. Urinary Si was analyzed by ICP-OES as previously described (24).

**Inter-Organism Homology of Si Transporters**

Homology search was performed with BLASTp (3) against plant and diatom sequences in the EMBL/GenBank nonredundant protein database using the default settings (http://www.ncbi.nlm.nih.gov/). BLASTp and tBLASTn were also used to identify homologs in a range of fully sequenced vertebrate genomes from the EMBL/GenBank and Ensembl databases, and also to identify homologs in selected phylogenetically relevant groups where complete genomes were not available (see Supplemental Table S1; Supplemental Material for this article is available at the Journal website). An alignment of homologs was generated using MUSCLE (http://www.ebi.ac.uk/Tools/msa/muscle/) under the default settings, producing a final alignment of 38 sequences from 17 species. ProtTest (1) found that the JTT + G + I model provided the best fit to the data under the Akaike Information Criterion. Maximum likelihood analysis was carried out using PhyML (19). Starting trees were generated by BioNJ, with tree searching using the NNI heuristic methods, and topology and branch lengths were optimized in ML calculations. One hundred bootstrap data sets were analyzed using the same model and method as for the PhyML analysis, with bootstrap proportions added as numbers to the nodes of the PhyML tree. The alignment was also used for Bayesian MCMC analysis using Phylobayes 3.3 software (26), under the CAT + G + I model until convergence (maximum discrepancy < 0.3, effective size > 100), for 15 parallel chains with sampling every 100 cycles and a burn-in equal to one-fifth the total size of the chain. Posterior probabilities were used to express the support for the nodes in the Bayesian phylogeny. The trees generated were viewed using FigTree (version 1.3.1, Andrew Rambaut, Institute of Evolutional Biology, University of Edinburgh 2006–2009).
Calculated Oxoacid Volumes

The structure for each oxoacid/oxoanion was optimized using the PB0 functional (2, 38, 39) and 6–31+g(d,p) atomic orbital basis set (4, 9, 15, 16, 20, 41, 42). The electron density corresponding to these optimized structures was used to estimate the molecular volume that describes the solvent accessible surface, defined as the volume bounded by a density contour of 0.001 electrons/Bohr³. An increased density of points was used to ensure a more accurate integration so that the computed molecular volumes are quantitative (37, 56). Since these species are in an aqueous environment, structures were optimized within a solvent field using the integral equation formalism variant of the polarizable continuum model (7, 52, 57) to account implicitly for the effects of an aqueous environment on the solvent accessible surface. The Gaussian09 suite of programs (17) was used in these determinations.

Transport Activity in Xenopus laevis Oocytes

Cloning the gene of interest and oocyte preparation. A cDNA sequence verified Rattus norvegicus IMAGE clone pExpress-1/Slc34a2 (Unigene ID: Rn.16933, Entrez Gene: 84395 in DH10BToNA) was purchased from Source BioScience LifeSciences (Cambridge, UK).

For synthesis of capped RNA, the open reading frame (ORF) was amplified by PCR with the following primers: 5' = GAGGATCCATG-GCTCCTTGGCCCGAGTTG-3' and 5' = GAGGATCCTAGAACACTGTAGTGTGGACA-3'. The fragment containing the ORF was inserted into the BglII site of a Xenopus oocyte expression vector, pXBG-ev1 (a pSTP64 T-derived pBluescript type vector into which Xenopus β-globin 5' and 3' UTR had been inserted) (40). Capped RNA was then synthesized from linearized pXBG-ev1 plasmids by in vitro transcription with mMESSAGE mMACHINE High Yield
Capped RNA Transcription Kit (Ambion) according to the manufacturer’s instructions.

Oocytes were isolated from *Xenopus laevis* frogs purchased from NASCO (Nasco-Fort Atkinson, WI) and from Watanabe Zosyoku (Hyogo Pref, Japan). Procedures for defolliculation, culture condition, and selection were the same as described previously (35). A volume of 50 nl of the in vitro cRNA transcripts (1 ng/ml) was injected into stage V oocytes using a Nanoject II automatic injector (Drummond Scientific). Water-injected oocytes were used as a negative control, *OsLsi1*-injected oocytes were used as positive controls while testing for influx activity, and *OsLsi2*-injected oocytes were used as positive controls while testing for efflux activity. Ethical approval was obtained (permit no. 21031043) from the Animal Care Committee of Laval University (Quebec City, QC, Canada).

**Influx transport activity.** After incubation in a Modified Barth’s Saline (MBS) solution (88 mM NaCl, 1 mM KCl, 2.4 mM NaHCO3, 15 mM Tris HCl at pH 7.6, 0.3 mM Ca(NO3)2, 0.41 mM CaCl2, 0.82 mM MgSO4, 10 μg/ml sodium penicillin, and 10 μg/ml streptomycin sulfate) at 18°C overnight, the cRNA-injected oocytes were exposed to the MBS solution supplemented with 1 mM H4GeO4, 0.1 mM HAAsO42- or 1 mM HPO42- at pH 7.6. Following 30 min incubation at 18°C, the oocytes were washed five times with MBS alone and digested with concentrated (61%) HNO3. The Ge, As, and P concentrations in the digested solutions were determined by ICP-MS (7700X: Agilent Technologies) with appropriate standards, QCs and sample blanks.

To investigate the Si influx and its dependence on extracellular [Na+] or [HPO42-], oocytes were incubated for three days at 18°C in MBS5 (84 mM Na+ and 2 mM HPO42-) supplemented with 100 μM each of penicillin and streptomycin. Then a set of 10 oocytes for each condition was exposed to MBS2 (1.7 mM H4SiO4, 10 mM Na+, and 0.5 mM HPHPO42-) or MBS3 (1.7 mM H4SiO4, 84 mM Na+, and 2 mM HPO42-) solution for 2 h. After exposure, oocytes were rinsed in a solution containing 0.32 M sucrose and 5 mM HEPES (pH 7.4) and then digested in 25 μl concentrated nitric acid, dried at 82°C for 2 h, reconstituted in plasma grade water (100 μl) and 10 μl analyzed by atomic absorption spectroscopy (see below).

**Efflux transport activity.** To investigate the efflux transport activity for H4GeO4 by *RnSlc34a2*, 50 nl 1 mM H4GeO4 in MBS were directly injected into *RnSlc34a2*-transfected oocytes. The oocytes were then washed five times with MBS and transferred to 200 μl of fresh MBS at 18°C. H4GeO4 was allowed to efflux into the incubation medium.

![Fig. 2. Correlation between renal *Rn*Slc34a2 expression (by quantitative RT-PCR analysis) and fasting urinary Si excretion. Urinary Si excretion in the rats (●, Si deplete; ○, Si replete) and laboratory chow reference group (▲, Si high reference) showed an inverse relationship with Slc34a2 expression in the kidneys; r = 0.47.](http://ajpcell.physiology.org/)

**Fig. 3. Immunohistochemistry analysis of Slc34a2 in freshly harvested rat kidney cortex.** Sections of freshly harvested kidneys from a healthy wild-type rat were analyzed by immunohistochemistry with anti-Slc34a2 (green) antibody (this figure) or the appropriate isotype control (data not shown). Cell nuclei were counterstained (blue) with Hoechst 33342 and cell cytoskeleton (f-actin, red) with phalloidin CF633. Antibody-stained sections and isotype controls were collected under identical settings, as stated in MATERIALS AND METHODS. A threshold removing all Slc34a2 attributable signal was defined on the isotype controls and uniformly applied to all images (i.e., antibody-stained images). Staining for Slc34a2 within the tubular epithelial cells was distinctly punctate so, as well as the signal above the isotype control being presented in an as-collected “intensity” format (i.e., the more secondary antibody that is bound, the brighter the signal) (A and B), it is also displayed as a binary format (i.e., all signal that is brighter than isotype threshold is given the maximum intensity value) as this aids visualization (C and D). All images are of the kidney cortex and scale bars are 50 μm. B: as-collected “intensity” format without actin staining. D: a high-power image (×63 magnification) of the area within the quadrant in image (C).
HEPES (pH 7.4), digested with concentrated HNO₃ (25oocytes were rinsed in a solution containing 0.32 M sucrose and 5 mM solution; Supplemental Table S2) for 0, 1, or 2 h. After exposure, concentrations of Na
concentrated HNO₃ and the samples were analyzed for Ge by ICP-MS and at the end of the experiment, the oocytes were digested with
and drinking water (Si-high reference; Fig. 4. Fasting urinary phosphorus excretion. Urinary P excretion was measured in the laboratory chow reference group (Si-high reference; Fig. 1A). The gene array findings1 were confirmed by real-time RT-PCR analysis, and this technique was also subsequently used to investigate a larger cohort of samples from the Si deplete (n = 10), Si replete (n = 8), and a reference group (n = 5) that were rats kept on a normal laboratory chow diet that is naturally high in Si (referred to as Si-high reference group). Slc34a2 (type II sodium-phosphate cotransporter), commonly referred to as NaPi-IIb, was expressed especially highly in the kidneys of rats on the Si deplete diet (2.8- and 4.8-fold higher than for kidneys from rats on the Si replete and Si-high reference diets, respectively; Fig. 1B). mRNA expression of other Slc34 family members, namely Slc34a1 and Slc34a3, were unchanged with dietary Si intervention (inset, Fig. 1B).

Correlation between Slc34a2 expression and urinary Si concentration showed an inverse exponential relationship between fasting urinary Si level and the relative expression of Slc34a2 (Fig. 2), implying that Slc34a2 is involved in the reabsorption of H₄SiO₄ from the pre-urine under dietary Si

<table>
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<th>Implicit Aqueous Solvent V₉ₕ, cm³/mol</th>
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1 Gene array data have been submitted to the Gene Expression Omnibus repository and assigned the reference GSE58404.
compared with the reference group (Fig. 4), showing that in the Si replete group) showed no difference in urinary P levels. However, the Si deplete group (with the same dietary P level as the Si-high reference group diet) showed significantly reduced P excretion (by 89 mg/mmol creatinine for the Si-deplete group) we measured urinary P excretion in the three groups. The Si-high reference group diet was higher in P than the Si-replete group diet, being 7.0 vs. 2.3 mg/g, respectively, and so, as expected, urinary P excretion was significantly reduced in the latter [by 89 mg/mmol creatinine for the means; \( P = 0.0001 \) in both instances]. Water-injected oocytes were used as a negative control. E: in H4SiO4 efflux studies, rice transporter Ls1 was used as a positive control. Data were corrected against water-injected control oocytes. F: changes in sodium and phosphate concentrations of the medium did not influence H4SiO4 influx by RnSlc34a2-expressing oocytes, nor that by OsLs1-expressing oocytes (\( P < 0.0001 \) in both instances). Water-injected oocytes were used as a negative control. E: in H4SiO4 efflux studies, rice transporter Ls1 was used as a positive control. Data were corrected against water-injected control oocytes. F: changes in sodium and phosphate concentrations of the medium did not influence H4SiO4 influx by RnSlc34a2-expressing oocytes. Data are shown as means \( \pm \) SE (\( n = 15 \)).

**Fig. 5.** Transport activity in RnSlc34a2-expressing oocytes. A–C: influx transport activity of Rattus norvegicus Slc34a2 for arsenate, HAsO\(_4\)\(^{2-}\) (\( P = 0.0001 \)) (A), phosphate, HPO\(_4\)\(^{2-}\) (\( P = 0.0008 \)) (B), and silicic acid, H\(_4\)SiO\(_4\) (\( P = 0.66 \)) (C). Rice transporter Ls1 was used as a positive control for H\(_4\)SiO\(_4\) influx (\( P < 0.0001 \)). D: the concentrations of sodium and phosphate in the medium did not influence H\(_4\)SiO\(_4\) influx by RnSlc34a2-expressing oocytes, nor that by OsLs1-expressing oocytes (\( P < 0.0001 \) in both instances). Water-injected oocytes were used as a negative control. E: in H\(_4\)SiO\(_4\) efflux studies, rice transporter Ls1 was used as a positive control. Data were corrected against water-injected control oocytes. F: changes in sodium and phosphate concentrations of the medium did not influence H\(_4\)SiO\(_4\) influx by RnSlc34a2-expressing oocytes. Data are shown as means \( \pm \) SE (\( n = 15 \)).

**Fig. 6.** Germanium transport activity in RnSlc34a2-expressing oocytes. A–F: uptake activity of Rattus norvegicus Slc34a2 for germanium, H\(_4\)GeO\(_4\) (\( P = 0.0001 \)) (A), phosphate, HPO\(_4\)\(^{2-}\) (\( P = 0.0004 \)) (B), and silicic acid, H\(_4\)SiO\(_4\) (\( P = 0.004 \)) (C). Rice transporter Ls1 was used as a positive control for H\(_4\)SiO\(_4\) influx (\( P < 0.0001 \)). D: the concentrations of sodium and phosphate in the medium did not influence H\(_4\)SiO\(_4\) influx by RnSlc34a2-expressing oocytes, nor that by OsLs1-expressing oocytes (\( P < 0.0001 \) in both instances). Water-injected oocytes were used as a negative control. E: in H\(_4\)SiO\(_4\) efflux studies, rice transporter Ls1 was used as a positive control. Data were corrected against water-injected control oocytes. F: changes in sodium and phosphate concentrations of the medium did not influence H\(_4\)SiO\(_4\) influx by RnSlc34a2-expressing oocytes. Data are shown as means \( \pm \) SE (\( n = 15 \)).

deprivation. No such relationship was observed for Slc34a1 and Slc34a3, or other candidate transporters identified in the gene arrays (Fig. 1A).

Only a few reports have demonstrated the renal expression of Slc34a2. The original paper characterizing the transporter demonstrated its presence in murine kidney at the mRNA level (23). Suyama et al. (50) confirmed this more recently by in situ hybridization as well as protein expression and localization by antibody staining. The kidney samples from our study were not adequately collected for immunohistochemical analysis, but were for RNA analysis. Thus we confirmed with appropriately collected kidneys from a control rat that, as previously published (50), Slc34a2 protein is expressed by the tubular epithelial cells of the kidney cortex (Fig. 3). Here, as previously reported (50), Slc34a2 showed distinct punctate staining: some of which was basolateral within the cell and some of which was apical/cytosolic (Fig. 3). Whether silicate deficiency dictates only the level of expression of Slc34a2 (Fig. 1) or, also its precise location in the cell, as excess dietary phosphate does (50), should be investigated in future work.

Finally, to translate these observations (i.e., that Slc34a2 has some basolateral expression in kidney cells and is upregulated in Si deplete diets) we measured urinary P excretion in the three groups. The Si-high reference group diet was higher in P than the Si replete group diet, being 7.0 vs. 2.3 mg/g, respectively, and so, as expected, urinary P excretion was significantly reduced in the latter [by 89 mg/mmol creatinine for the medians; \( P = 0.008; n = 6 \) and 5, respectively (Fig. 4)]. However, the Si deplete group (with the same dietary P level as the Si replete group) showed no difference in urinary P levels compared with the reference group (Fig. 4), showing that in this group, phosphate was being (relatively) wasted as a consequence of Si being conserved.

**RnSlc34a2 Transport Activity**

The ubiquitous nature of Si makes transport studies of soluble silicic acid (H\(_4\)SiO\(_4\)) challenging. It is well known that related oxoacids may ride the same transport systems (22, 29, 36, 54) owing to similarities in their structure and solvated molecular volume (Table 1). Germanic acid (H\(_4\)GeO\(_4\)), the
closest structural analog of silicic acid, is therefore often employed as a proxy for Si transport, thereby avoiding all background and contamination issues with Si and facilitating analysis (22, 29). Recently, however, graphite furnace atomic absorption spectrometry (GFAAS) was shown to be effective for directly measuring Si influx/efflux in *Xenopus laevis* oocytes transfected with Si-transporting aquaporins from plants (8, 18). Hence, both methods of characterizing Si transport—indirect and direct—were used in the present investigation.

The Slc34a2 coding sequence was inserted into a *Xenopus laevis* expression vector, and cRNA synthesized from this construct was injected into oocytes. Initial expression and plasma membrane localization were verified using an eGFP-tagged Slc34a2 construct. Slc34a2 is recognized as a sodium phosphate importer, especially in the brush border of small intestine membrane cells (14, 31, 36, 55), and arsenate also rides this transport system (36, 54). Therefore, both of these oxoacids were utilized as easily measured probes to confirm Slc34a2 influx activity (Fig. 5, *A* and *B*). The rice Si importer *OsLsi1* was used as a positive control and was found to promote both H4SiO4 (Fig. 5 *C*) and H4GeO4 (Fig. 6 *A*) influx.

By contrast, no influx of either H4SiO4 or H4GeO4 by Slc34a2-expressing oocytes was observed (Figs. 5 *C* and 6 *A*).

The transmembrane domains were predicted using SOSUI software. The sequence highlighted in yellow through multiple sequence alignment of the three *Rattus norvegicus* Slc34 family members (Fig. 1 *C*) is present in the ninth transmembrane helix of Slc34a2 (shown in bold). The COOH-terminal and NH2-terminal amino acids for each transmembrane domain are indicated, as is the type of α-helical structure (i.e., primary or secondary helices, denoted as 1° and 2°, respectively).

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</tbody>
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The transmembrane domains were predicted using SOSUI software. The sequence highlighted in yellow through multiple sequence alignment of the three *Rattus norvegicus* Slc34 family members (Fig. 1 *C*) is present in the ninth transmembrane helix of Slc34a2 (shown in bold). The COOH-terminal and NH2-terminal amino acids for each transmembrane domain are indicated, as is the type of α-helical structure (i.e., primary or secondary helices, denoted as 1° and 2°, respectively).
Fig. 8. Phylogeny of SLC34a gene family member in vertebrates. A: the tree was produced using PhyML maximum likelihood analysis with the JTT + G + I model from an alignment of 880 positions. Numbers at nodes are a percentage of 100 bootstrap replicates, with nodes having <70% bootstrap support being collapsed. The tree was rooted using the single SLC34a homolog identified from the lamprey genome. The Slc34a gene phylogeny largely agrees with the species phylogeny for vertebrates (33), with incongruent branches (e.g., the basal branches of the a2 clade) only having low statistical support. The maximum likelihood phylogenetic analyses resolve that the Slc34a clade evolved from a single ancestor in jawless vertebrates, and likely involved two main duplication events, initially producing the a3 and a1 clades, with a further divergence of the a1 and a2 clades. A teleost-specific duplication event resulted in the evolution of Slc34a2a and Slc34a2b, as found in stickleback and zebrafish. The Bayesian analysis had poor phylogenetic resolution at the base of the a2 clade, but still resolves the a1 and a3 groups as distinct monophyletic clades, and is not incongruous with the maximum likelihood analysis results. For full details of the species and sequences used see Supplemental Table S1.

B: the tree was produced using Phylobayes Bayesian MCMC analysis under the CAT + G + I model (15 parallel chains with sampling every 100 cycles, burn-in one-fifth the total size of the chain) from an alignment of 880 positions. Numbers at nodes indicate posterior probabilities, with nodes having <0.95 support being collapsed. The scale bar indicates the average number of amino acid substitutions per site. The Slc34a1 clade is in green, the Slc34a2 clade is in blue, and the Slc34a3 clade is in red.
other hand, efflux of both H$_4$SiO$_4$ and H$_4$GeO$_4$ was detected for oocytes expressing Slc34a2 (Figs. 5E and 6B, respectively) as well as those expressing rice Si exporter OsLsi2, which was employed as a positive efflux control. Of note, the magnitude of fractional H$_4$SiO$_4$ efflux after 2 h was quantitatively similar between Slc34a2 and OsLsi2.

Given that inward phosphate (HPO$_4^{2-}$) transport by Slc34a2 is coupled to the inward transport of three sodium ions [i.e., it is electrogenic (14, 55)], we investigated how varying the concentrations of Na$^+$ and HPO$_4^{2-}$ in the external medium might influence Si influx and efflux in Slc34a2-expressing oocytes. No significant effects were observed at the broad concentrations investigated (Figs. 5D and F). These findings suggest that Si is not translocated across the membrane through the Na$^+$ or HPO$_4^{2-}$ transport site, but through an independent transport site that is potentially involved in Si efflux primarily. In keeping with this possibility is the presence of multiple, often independent binding sites in a number of ABCD family members (47). Alternatively, Slc34a2 could cooperate with accessory proteins to promote Si efflux. In this regard, the Na$^+$/K$^+$-ATPase $\gamma$-subunit FXYD2 appears to play a role in Mg$^{2+}$ transport, while the $\alpha$- and $\beta$-subunits alone do not exhibit such transport capabilities (46).

Homology Between Slc34a2 and Si Transporters

Comparative sequence analysis of Slc34a2 indicated no significant homology with known plant or algal Si transporters. However, marked similarities were revealed upon pairwise alignment of the transmembrane domains of Slc34a2 and the plant Si efflux transporter Lsi2 (Fig. 7), thereby suggesting a conserved structure among Si efflux proteins.

Phylogenetic Analysis of the Slc34 Family

Sequence alignment within the rat Slc34 gene family led to the identification of a ~30-residue stretch that is only present in Slc34a2. Given that Slc34a1 and Slc34a3 were not upregulated under Si deprivation, this finding points towards the possibility that the ~30-residue stretch conveys Si transport activity to Slc34a2 (Fig. 1C and Table 2).

Phylogenetic analysis of the Slc34a genes from a range of vertebrates (Supplemental Table S1 and Fig. 8) showed that the family underwent an expansion relatively early in vertebrate evolution, resulting in three distinct main groups (a1, a2, and a3) among the modern jawed vertebrates. At least one member of the a2-group was found in all of the jawed vertebrate genomes searched. In contrast, losses of the a1- and a3-group
genes were observed in several fully sequenced genomes (e.g., zebrafish, chicken). This would suggest that the Slc34a2-group genes have a unique or important role whose loss cannot be complemented for by other transporters, and that this function is conserved across the jawed vertebrates. Common to all members of the Slc34a2 group, and to the homologous Slc34a2 gene in the lamprey, is a motif containing three positive amino acid residues (R, H, or K) separated by smaller uncharged residues (commonly C or S) (Fig. 9). This motif aligns with the unique predicted transmembrane domain noted above in the rat Slc34a2 gene (Fig. 1C and Table 2), and points towards an important functional role. Conserved positively charged amino acids have been noted in other Si-related proteins, such as the GRQ motifs of the SIT active Si transporters (32, 31). It may be postulated that these residues interact with local negative charges on the silicic acid molecule as part of a general biochemical basis for transmembrane Si transport.

Conclusions

The identification that Slc34a2 can transport Si in mammals establishes another distinct gene family of Si transporters that could be involved in the regulation of Si homeostasis and that bears no sequence similarity with known Si-related genes in plants, sponges, choanoflagellates, or diatoms, although it shows strong structural similarities to silicon exporters in plants. Crucially, our work is also one of the first pieces of evidence for a functionally relevant Si-responsive gene in mammals. In parallel with this work, Carneau et al. (18) and Deshmukh et al. (8) have recently identified Si-permeable aquaporins that appear to play an important role in Si influx. Coupled with the active efflux transporter that is reported herein, we propose a Si transport model in mammals that mirrors that known in plants (28), i.e., a model in which an influx and efflux transporter must be present to allow Si movement through cells. Here, Slc34a2 is effluxing H$_2$SiO$_4$ from the renal tubular epithelial cell into the circulation, i.e., it is involved in the reabsorption of H$_2$SiO$_4$ in the kidneys. As an inevitable consequence of this expression at this cellular location, phosphate will be moved in the opposing direction. Dietary Si-P interactions have been noted, with animals on a Si-deficient diet showing conserved bone Si levels but depleted bone P levels (24). Assuming Slc34a2 is similarly involved in bone conservation of Si as it is in the kidney then our results explain these observation (24). Finally, it is also interesting to note that Lsi2’s are equally upregulated in plants in conditions of Si deprivation (34), a phenomenon that was instrumental in identifying Slc34a2 in this work. Collectively, our data provide indication that, rather than being a biochemically inert element, Si in fact plays a role in vertebrate physiology deserving of its preservation under exposure conditions of deprivation.

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DISCLOSURES

No conflicts of interest, financial or otherwise, are declared by the author(s).

AUTHOR CONTRIBUTIONS


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IDENTIFICATION OF A MAMMALIAN SILICON TRANSPORTER


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