

Disruption of urate transport in familial renal glucosuria and report on SGLT2 expression in normal and pathological kidney

Alteração do transporte de ácido úrico na Glicosúria Renal Familiar e expressão de SGLT2 no rim normal e patológico

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ABSTRACT

Familial renal glucosuria (FRG) is a rare co-dominantly inherited benign phenotype characterized by the presence of glucose in the urine. It is caused by mutations in the *SLC5A2* gene that encodes SGLT2, a Na⁺-glucose co-transporter. The purpose of our current work was twofold: to characterize the molecular and phenotype findings of an FRG cohort and, in addition, to detail the SGLT2 expression in the adult human kidney. The phenotype of FRG pedigrees was evaluated using direct sequencing for the identification of sequence variations in the *SLC5A2* gene. The expression of SGLT2 in the adult human kidney was studied by immunofluorescence on kidney biopsy specimens. In the absence of renal biopsies from FRG individuals, and in order to evaluate the potential disruption of SGLT2 expression in a glucosuric nephropathy, we have selected cases of nucleoside analogues induced proximal tubular toxicity. We identified six *novel SLC5A2* mutations in six FRG pedigrees and described the occurrence of hyperuricosuria associated with hypouricaemia in the two probands with the most severe phenotypes. Histopathological studies proved that SGLT2 is localized to the brush-border of the proximal tubular epithelia cell and that this normal pattern was found to be disrupted in cases of nucleoside analogues induced tubulopathy. We present six *novel SLC5A2* mutations, further contributing to the allelic heterogeneity in FRG, and identified hyperuricosuria and hypouricaemia as part of the FRG phenotype. SGLT2 is localized to the brush-border of the proximal tubule in the adult human normal kidney, and aberrant expression of the co-transporter may underlie the glucosuria seen with the use of nucleoside analogues.

Key words: glucosuria; proximal tubule; SGLT2 expression; uric acid.

■ RESUMO

A Glicosúria renal familiar (GRF) é um fenótipo de transmissão autossômica co-dominante, causada por mutações do gene *SLC5A2* que codifica SGLT2, o principal co-transportador de Na⁺-glucose do túbulo proximal. Caracteriza-se pela redução da reabsorção tubular renal de glucose, causando glicosúria na ausência de hiperglicemia e de outros sinais de disfunção tubular. O objectivo do trabalho aqui apresentado foi caracterizar a nível molecular e fenotípico um conjunto de famílias afectadas por esta entidade e ainda detalhar a expressão de SGLT2 “in vivo” em amostras de tecido renal normal e patológico. Avaliámos fenotipicamente um grupo de indivíduos afectados por GRF e identificámos por sequenciação directa as alterações encontradas ao gene *SLC5A2*. A expressão de SGLT2 no rim adulto humano foi estudada por microscopia de fluorescência. Na ausência de biópsias renais de indivíduos afectados com GRF e de modo a avaliar potenciais alterações de expressão de SGLT2 em nefropatias com glicosúria, seleccionamos casos de lesão tubular proximal por toxicidade de análogos de nucleósidos. Identificamos seis novas mutações de *SLC5A2* em indivíduos de seis famílias. Nos indivíduos com glicosúria mais severa observamos igualmente hiperuricosúria com hipouricemia. Pudemos demonstrar que no rim adulto normal, SGLT2 está presente na bordadura em escova do epitélio do tubo proximal e que em casos de tubulopatia dos análogos dos nucleósidos a sua expressão está diminuída. Apresentamos assim seis novas mutações de *SLC5A2* que contribuem para a heterogeneidade alélica da GRF e identificámos a hiperuricosúria com hipouricemia como características deste fenótipo. Confirmámos a expressão de SGLT2 na bordadura em escova do túbulo proximal e demonstramos que a diminuição da sua expressão pode estar subjacente à glicosúria observada na terapêutica com análogos dos nucleósidos.

Palavras chave: ácido úrico; expressão de SGLT2; glicosúria; tubo proximal.

■ INTRODUCTION

Familial renal glucosuria (FRG) is characterized by the presence of glucose in the urine in the absence of diabetes *mellitus* or generalized proximal tubular dysfunction. Mutations in the *SLC5A2* gene, positioned in 16p11.2 and encoding SGLT2, the Na⁺-coupled glucose transporter that accounts for the bulk of glucose reabsorption in the renal proximal tubule, are responsible for the large majority of FRG pedigrees¹. Most of these mutations are private and usually involve *missense* alleles, although *nonsense*, small deletions (both *in-frame* and *frameshift*) and splice site mutations have been reported. Even though an increasing number of FRG cohorts is being reported²⁻⁵, patients presenting with severe forms of FRG with glucosuria in excess of 50 g/1.73m²/24h are distinctly rare.

Two members of the Na⁺-coupled glucose transporter family, SGLT2 and (to a lesser extent) SGLT1, are responsible for the secondary active uptake of glucose from the proximal lumen at the apical side

of the epithelia. Once within the cell, glucose is returned to the systemic circulation by facilitative transport mediated by GLUT1 and 2⁶.

Renal tubular glucose transport has been pointed out as a target in the treatment of type 2 diabetes *mellitus* (T2DM) and several SGLT2 inhibitors (SGLT2i) are now awaiting FDA approval. In a previous report, we anticipated that these compounds were likely to have a diuretic effect, based on the findings of moderate volume contraction with secondary activation of the renin-angiotensin-aldosterone system in severe FRG phenotypes^{2,3}. This was later ascertained in clinical trials with SGLT2i^{7,8}

Our group performs routine molecular characterization of FRG pedigrees and, whenever possible, phenotypic evaluation of such individuals. We now report the molecular and clinical findings of six pedigrees displaying *novel SLC5A2* alleles. For two of these probands, glucosuria was in the range of the urinary glucose excretion (UGE) induced by pharmacological inhibition of SGLT2. One interesting and consistent

observation with the administration of SGLT2i, is the almost dose-proportional decrease in serum uric acid levels^{7,8}, which is paradoxical in light of the previously mentioned diuretic effect. We have, therefore, evaluated severe FRG phenotypes regarding serum and urinary uric acid levels. In addition, we assessed SGLT2 protein expression in adult normal human kidney and validated an FITC-immunofluorescence technique for paraffin-embedded kidney biopsy samples, with the purpose of detailing the topological location of SGLT2. Both techniques were performed using an available commercial antibody. In the absence of renal specimens from FRG individuals, we have searched for aberrant SGLT2 expression in renal biopsies of patients with nucleoside analogues induced proximal tubular toxicity and glucosuria.

SUBJECTS AND METHODS

Phenotype evaluation and mutation analysis of FRG pedigrees

This study was part of the medical evaluation for glucosuria. After obtaining informed consent from participating individuals or their legal guardians, individuals were enrolled. Mutation analysis was performed as previously reported⁹, but for exons 10 to 12, a single amplicon of 978 bp was PCR amplified and directly sequenced, using the following forward 5'-CTCGTCGTGAAGCTCATGC-3' and reverse 5'-CCAACCCTCAGTCGAGAAAT-3' primers. Only individuals bearing *SLC5A2* mutations not previously described were included in this report.

One hundred control chromosomes were assayed for the presence of the *missense* mutations identified, excluding them as common polymorphisms. We used direct sequencing of exons 5 and 12, for the c.571 A>C and c.1475 G>T alleles, respectively. In the case of the c.1343 A>T transversion, a new *TspRI* restriction site was inserted within the 978 bp exon 10-12 amplicon, therefore enabling easy testing by PCR-RFLP of affected members and control chromosomes.

Immunohistology of kidney biopsies

SGLT2 detection by immunofluorescence was performed in five formalin-fixed and paraffin-embedded

kidney biopsies with no detectable histopathological changes. Two additional sample of known proximal tubular toxicity with glucosuria induced by tenofovir or adefovir were also included. Heat-induced epitope retrieval was applied to 3µm sections with Target Retrieval Solution pH 9 (ref. S2367, Dako) at 95-99°C for 20 min. Incubations with primary goat polyclonal antibody anti-human SGLT2 C-terminal domain, from residues 590-640 (ref sc-47401, from SantaCruz Biotechnology (1:5) and a secondary antibody donkey anti-goat IgG-FITC (1:20) – ref sc-2420, Santa Cruz Biotechnology – were carried out for 60` at room temperature.

RESULTS

Phenotype evaluation and mutational analysis

We identified six *novel* alleles in 13 individuals, from six different pedigrees. The available clinical data is detailed in Table I. Relatives were screened for glucosuria by dipstick urine analysis and, whenever possible, quantified by a 24h urine collection. All affected individuals had normal renal function. Two heterozygous, one compound heterozygous and three homozygous probands were identified. The two probands with the most severe phenotypes, with UGE of 87.8 and 43.7 g/1.73m²/24h – individuals I.1 of pedigree 1 and II.1 of pedigree 2 – also had hyperuricosuria: excreting 1242 and 990 mg/1.73m²/24h (reference values 250-750), with an urinary uric acid (mg)/Kg of body weight of 19.1 and 21.5 (reference values in a healthy population: 7.6±3.75 [adolescents] and 7±1.6 [adults]¹⁰, respectively. For proband of pedigree 1 (the most severe of our cohort) this was clearly associated with hypouricaemia of 1.9 mg/dl (reference range: 2.5-7). Proband II.1 of family 2, although having a serum uric acid of 3.5 mg/dl at the time of evaluation, by the age of eight years the reported value was of 2.4 mg/dl.

Three mutated alleles are *missense* substitutions: p.T191P, p.Q448L and p.G492V. All represent non-conservative substitutions (with the exception of Val for Gly, in that both are non-polar aminoacids) affecting highly conserved residues among SGLT members (Fig. 1). Their putative pathogenicity was further assessed *in silico* using the bioinformatic algorithm *Polyphen* and, as such, all were predicted

Table 1

Phenotype evaluation and mutational analysis in affected individuals and their relatives.

Family members (age*)	Country of origin	Weight (Kg)	Height (cm)	Glucose excretion (g/1.73m ² /24h)	Plasma glucose (mg/dl)	HbA1c (%)	Serum uric acid (mg/dl)	Urinary uric acid		allele 1 (cDNA;protein)	allele 2 (cDNA;protein)
								mg/1.73m ² /24h	mg/Kg		
Family 1 I.1 (35)	Portugal	65	167	87.8	74	5.4	1.9	1242	19.1	c.1030_1057del;p.V346fsX17	c.1030_1057del;p.V346fsX17
Family 2 I.1 (na)	Costa Rica	na	na	4+	na	na	na	na	na	c.1946 G>A;p.W649X	wt
I.2 (na)		na	na	1+	na	na	na	na	na	wt	wt
II.1 (10)		25	126	43.7	96	5.9	3.5	990	21.5	c.1946 G>A;p.W649X	wt
Family 3 I.1 (35)	Turkey	na	na	14.2	na	na	na	na	na	IVS7+5g>a	c.1343A>T;p.Q448L
I.2 (30)		na	na	5.6	na	na	na	na	na	c.1343A>T;p.Q448L	wt
II.1 (18)		60	170	9.8	92	4.8	5.9	na	na	c.1343A>T;p.Q448L	c.1343A>T;p.Q448L
II.2 (14)		49	152	30	87	4.2	5.3	na	na	IVS7+5g>a	c.1343A>T;p.Q448L
II.3 (8)		21	117	13	84	5.5	2.2	na	na	c.1343A>T;p.Q448L	c.1343A>T;p.Q448L
Family 4 I.1 (10)	Turkey	31	140	25	90	5.5	4.4	na	na	c.1566C>A;p.C522X	c.1566C>A;p.C522X
Family 5 II.1 (16)	Turkey	58	155	14.4	71	4.1	3.4	na	na	c.571A>C;p.T191P	c.571A>C;p.T191P
II.2 (11)		32	139	13.2	87	4	4.1	na	na	c.571A>C;p.T191P	c.571A>C;p.T191P
Family 6 I.1 (53)	Portugal	82	174	4.5	114	5.4	3.6	na	na	c.1475G>T;p.G492V	wt

to be probably damaging. In addition, these sequence variations were not detected in 100 control chromosomes and were, therefore, excluded as common polymorphisms. There are three truncating mutations, including two *non-sense*, p.C522X and p.W649X and one *frameshift*, p.V346fsX17.

Family 3 displays a unique genetic feature. The proband, II.2, is a 14-year-old girl born of a consanguineous marriage. Yet, her severe phenotype is not related to consanguinity but to compound heterozygosity: in addition to the ancestral p.Q448L mutation, shared by both parents, she displays the donor splice

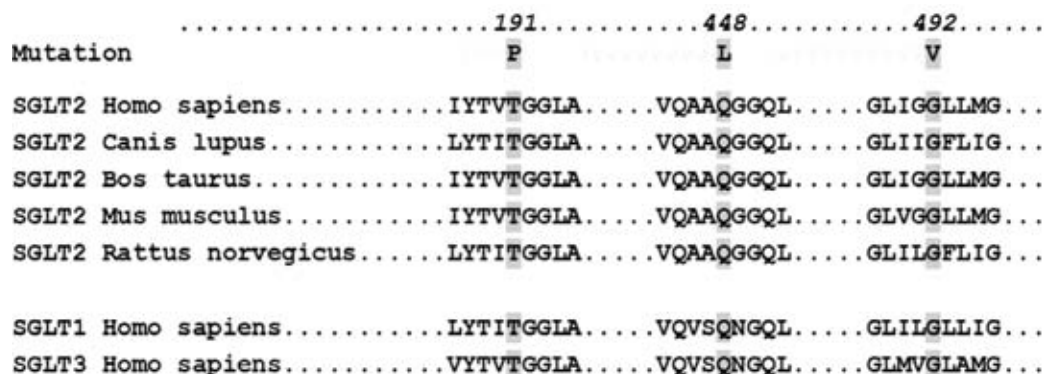


Figure 1

Sequence alignments: Multiple alignments of SGLT2 proteins from different species and comparison to human SGLT1 and SGLT3. The missense alleles of the current cohort are represented on top. The highly conserved residues are highlighted in grey

site mutation IVS7+5g>a transmitted by the father, in *trans* with the former allele. The remaining siblings are homozygous for the ancestral p.Q448L mutation, as anticipated in consanguineous marriages.

■ SGLT2 expression in human kidney

Immunofluorescence experiments in five kidney biopsies with no detectable histopathological changes localized SGLT2 to the brush-border of proximal

tubular epithelia in normal kidney (Fig. 2A). Two patients with nucleoside analogues induced tubulopathy and glucosuria were identified in our pathological database. The first patient presented with acute renal failure after the introduction of tenofovir for his HIV condition. At renal biopsy, serum creatinine (Scr) was 7.0 mg/dl (on dialysis) and the 24h UGE of 4.9 g (175 mg/dl). The second individual had been for several months on adefovir for hepatitis B infection, before referral to our department. At evaluation, Scr was 1.7 mg/dl and the 24h UGE of 15.4 g (358 mg/dl). By

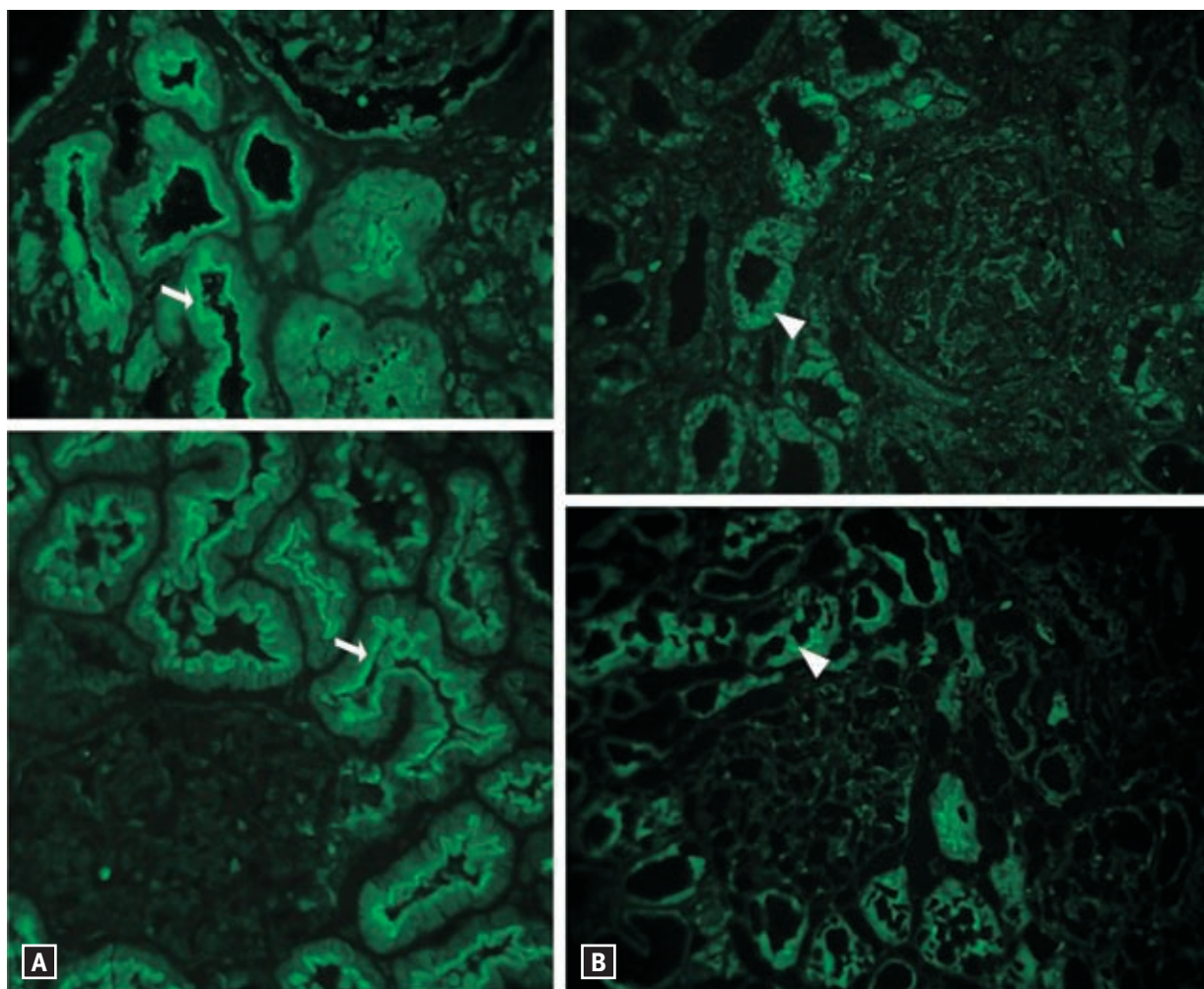


Figure 2

SGLT2 expression in human kidney. A) Immunofluorescence clearly localizes SGLT2 expression (white arrow) to the brush-border of proximal tubular cells in normal kidney(x400). B) Renal biopsies from patients with nucleoside analogues induced proximal tubular toxicity. Top: tenofovir case with loss of SGLT2 expression from the apical side of proximal epithelia (arrowhead)(x400). Bottom: adefovir case with significant reduction in the expression of the cotransporter, although some discontinuous pattern is still seen (arrowhead)(x400).

immunofluorescence, apical SGLT2 expression was abolished in the biopsy of first patient, while in the second case it was significantly reduced (Fig. 2B).

■ DISCUSSION

The kidney contributes to glucose homeostasis by reabsorbing at the proximal tubule virtually all of the glucose filtered by the glomerulus, in a two-step process. Firstly, glucose and Na⁺ are sequentially cotransported from the tubular lumen via secondary active transport mediated by the renal Na⁺-glucose co-transporters SGLT2 and SGLT1. Secondly, glucose is released from tubular epithelial cells into the systemic circulation by facilitative Na⁺-independent glucose transporters GLUT1 and GLUT2. SGLT2 is a low-affinity, high-capacity transporter, predominately expressed at the most proximal S1 tubular segment of the nephron¹¹. Under physiological conditions, SGLT2 reabsorbs the majority of the filtered glucose¹² and is considered a candidate gene for FRG. Mutations in SGLT1 account for the glucose-galactose malabsorption syndrome (GGM), which is characterized by severe diarrhoea upon the introduction of glucose or galactose to the diet of infants¹³. The different phenotypes of FRG and GGM reflect the differential expression pattern of both co-transporters as assessed by RT-PCR: SGLT1 being expressed mainly in intestinal epithelia, while SGLT2 is mostly restricted to the proximal renal tubule¹⁴. Another important difference between both cotransporters is the apparent difficulty in assessing SGLT2 expression by immunodetection, as compared to SGLT1. In fact, most of the work for SGLT2 was done by *in-situ* hybridization or RT-PCR^{11,14}.

This report aims at further clarifying the renal tubular glucose transport by reporting the phenotype and genotype findings of FRG pedigrees displaying *novel* SGLT2 alleles, particularly for those individuals with severe forms, and also by assessing SGLT2 protein expression in adult human kidney.

For the six pedigrees analysed, there are six *novel* *SLC5A2* alleles and the *IVS7+5g>a* mutation already reported. According to the previously described co-dominant pattern of inheritance for FRG, those individuals homozygous or compound heterozygous for a given mutation were shown to have larger UGE, when compared to heterozygous patients. For this

cohort, the previously defined 10 g/1.73m²/24h cut-off appropriately seems to apply, with the exception of proband from pedigree 2. In this individual having an unusual severe phenotype, only one mutation was characterized. In fact, a second sequence variation in intron 3 (*IVS3-20g>a* [data not shown]) was identified, but after testing the remaining family members, it was found to be in *cis* with the *nonsense* mutation. It is possible that the second allele might have been entirely missed, either because it resides outside the coding region or it consists on a large DNA rearrangement not amenable to detection by routine PCR based techniques. Moreover, a dominant negative effect or transheterozygosity with another gene coding for a protein also having a role in renal glucose transport, are alternative explanations.

All characterized sequence variations are expected to be deleterious for SGLT2 function: the three *missense* mutations are predicted to be pathogenic substitutions, while the p.V346fsX17, p.C522X and p.W649X mutations, expected to lead to truncated proteins at SGLT2 transmembrane domains 3, 13 and 14, respectively. The *IVS7+5g>a* allele of Family 3 is recurrently reported in unrelated pedigrees of different ethnic origins¹⁵. Considering the significant allelic heterogeneity in FRG, with most mutations being family specific, the high recurrence for this single allele can be due to a mutational hot-spot. This pedigree represents the first instance of allelic heterogeneity in a consanguineous FRG pedigree, in that the proband is a compound heterozygous for the ancestral p.Q448L mutation, shared by both parents, and the splice site mutation in intron 7. This family furthermore illustrates the co-dominant nature of FRG, with all members being affected irrespectively of heterozygosity (mother), compound heterozygosity (father and proband) or homozygosity (remaining siblings) for the identified mutations.

Hyperuricosuria with hypouricemia is the most interesting phenotype finding in the probands presenting with the highest UGE, suggesting urinary urate over-excretion as the mechanism underlying hypouricaemia in FRG and in pharmacological SGLT2 inhibition^{7,8}. These observations also rule out the possibility of non-specific pharmacological inhibition of renal urate transporters like GLUT9, the facilitative glucose transporter encoded by the *SLC2A9* gene, mutated in familial renal hypouricemia¹⁶ and expressed at the proximal renal tubule in humans.

Interestingly, GLUT9 was also shown to exchange extracellular glucose for intracellular urate¹⁷. Two isoforms, GLUT9S (expressed at the apical side) and GLUT9L (at the baso-lateral side) act concertedly to reabsorb uric acid from the glomerular filtrate. However, in case of a high glucose concentration in the proximal tubular lumen, as seen in FRG and with SGLT2 pharmacological inhibition, the direction of urate transport could be reversed, with glucose being exchanged for urate and leading to excessive urinary excretion and secondary hypouricaemia.

We have detailed the expression of SGLT2 in human adult kidney by assessing its expression by immunofluorescence in biopsy specimens. A previous report⁴ had already localized SGLT2 to the apical side of proximal renal tubular epithelia in humans. Our experiments confirm that SGLT2 is, in fact, expressed at apical side of normal proximal tubular epithelia and confined to the brush-border. Importantly, its expression is distinctly reduced or absent from the apical side of the proximal tubule in the cases of antiretroviral renal toxicity, both presenting with proximal tubulopathy with glucosuria. These anticipated “negative controls” not only confirm the specificity of the antibody used, but also identify the loss for SGLT2 expression as the mechanism underlying the glucosuria induced by some nucleoside analogues.

In conclusion, we present the phenotype and genotype characterization of six FRG pedigrees displaying six *novel* SGLT2 mutations. The hyperuricosuria found in FRG is a plausible direct cause for the hypouricaemia in the setting of severe glucosuria, either occurring naturally -as in FRG- or induced by pharmacological inhibition of SGLT2. We theorized about GLUT9 being the transporter responsible for these observations, by means of an antiporter mechanism. Finally, we localized SGLT2 in the human adult normal kidney, to the brush-border of proximal tubular epithelia cell.

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Conflict of interest statement: None declared.

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