Glucose transporters in kidney; the role of gender and diabetes mellitus

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Introduction
Facilitative D-glucose transport is mediated by members of the solute carrier transporter, glucose transporters (GLUTs), or uniporters family. Sodium-glucose co-transporters (SGLTs) involve in the D-glucose co-transport along with sodium, and its types 1and 2 (SGLT1 and SGLT2) are proteins that are encoded by the solute carrier family 5 member 1&2 (SLC5A1 and and SLC5A2) genes, respectively. SGLT2 (a high-capacity carrier) expresses in the S1 and S2 segments, while SGLT1 expresses in the S2 and S3 segments in human, rat and mice (1,2). In normal condition, SGLT2 has the ability to reabsorb 95% or more of the filtered glucose, however, SGLT1 reabsorbs about 3% of the remained filtered glucose (3). Functional characteristics of SGLT2 and SGLT1 are coupling of Na+ glucose coupling ratios (1:1 and 2:1), therefore this allows transporting the small amounts of glucose to the late proximal tubule.

The mediate actions of SGLT1 and SGLT2 are also different. Principally, the SGLT2 mediates the reabsorption of glucose, while SGLT1 is involved more in glucose reabsorption in the small intestine where SGLT2 is not expressed. The glucose that not reabsorbed in the early section of tubule could be reabsorbed in the late section of the tubule by SGLT1.(1,3). Accordingly, glucose reabsorption by inhibition of SGLT1 in the kidney may perform a treatment process against the hyperglycemic condition, and on the other side, SGLT1 inhibitor alone or with SGLT2 as a therapeutic may use, for the treatment of hyperglycemia and diabetes mellitus.

Sex hormones, gender, and glucose transporters
A kind of dimorphism is involved in the glucose uptake regulation by sexual hormones. In the polycystic ovarian syndrome, GLUT expression and glucose metabolism are defected, and insulin resistance is occurred (4).

Implication for health policy/practice/research/medical education:
It seems that the co-transporters of sodium-glucose (SGLTs) function is gender related, and to use SGLTs inhibitors as therapeutic agents in diabetic condition, the role of sex needs to be more specified.

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Testosterone therapy in male and female animals decreases glucose transport by down-regulating of GLUT1 (5) while estradiol via receptor β shows a causing or producing diabetes (diabetogenic) activity (6). In physiological concentration, both sexual hormones; testosterone and estradiol, preserve a normal insulin sensitivity (4). Testosterone also may increase insulin sensitivity in diabetes type 2- induced hypogonadism (7). The higher kidney SGLT1 (rSGLT1) and SGLT2 (rSGLT2) expressions have been found in female rats compared with male rats, while the higher expression of SGLT2 has been detected in the kidney of male mice (2, 8,9).

The expression levels of SGLT1 and SGLT2 are reported to be sex-independent in human (10). It is reported that SGLT1 in the rat's kidney was a 75-kDa protein expressed mainly in the S3 segments of proximal tubule, and its expression was androgen and sex dependent (9). It is also reported that the amount of SGLT2 protein was male dominant in the mouse kidneys, while the SGLT2 mRNA expression was female dominant, and in rodents the SGLT2 expression was kidney-specific (8). The GLUT carriers and SGLTs are encoded by solute carrier family 2 (SLC2) and SLC5 genes respectively, and the main substrate of facilitative GLUT carriers are glucose, galactose, mannose, glucosamine, and the main substrate of SGLTs is biotin, lipoate, and pantothenate that all are substrates of GLUT1 (5) while estradiol via receptor β shows a causing or producing diabetes (diabetogenic) activity (6).

Glucose transporters and diabetes

In order to control the blood glucose in diabetic patients, it is necessary to reduce the development of metabolic abnormalities and decrease the risk of complications such as nephropathy and cardiovascular disorders. Accordingly, special attention has been made to SGLTs due to their role of kidney SGLT1 in diabetes and kidney damage (11). The relating mRNAs of the renal GLUT1 and SGLT isoforms were expressed at greater levels in female compared with male mice while no considerable sex difference was detected in serum level of glucose (12). Madunić et al reported that expression of SGLT1 in male is more than female mice and there is a female-dominant expression in rats (13). Moreover, it is clear that the expression of SGLT2 mRNA in female kidney was higher than male, but the SGLT2 protein expression was reversed, which reveals that the gender difference in renal SGLT2 protein is not explained by the expression of different levels of mRNA (8).

Uncontrolled hyperglycemia can overcome the delivery capacity of early proximal SGLT2 so increasing the glucose transport to the late proximal tubule occurs, which enhances the glucose reabsorption by SGLT1. Experimental research on animal diabetic model demonstrates a rise of renal protein expression of SGLT2, which contributed to increase renal glucose reabsorption capacity (16). In the renal cortex of streptozotocin-induced diabetic rats, an increase in mRNA expression for SGLT1 (17). Similarly, in diabetic obese Zucker rats, the level of renal SGLT1 mRNA was more than that of immature non-diabetic lean rats (18), and the smaller expression of SGLT1 mRNA but higher expression of renal membrane SGLT1 protein were detected in BTBR ob/ob T2DM mice (animal model of diabetic nephropathy) (19). The renal expression and activity of SGLT1 decreased in streptozotocin-induced diabetes (20). Additionally, the SGLT2 knockout mice showed a decreased expression of renal SGLT1 protein (21). Therefore, down-regulation in renal SGLT1 expression may limit glucose uptake and toxicity in the late proximal tubule when glucose load and reabsorption are enhanced (1, 21) while in normoglycemic mice, SGLT2 and SGLT1 are responsible for all renal glucose reabsorption (3). It is unclear whether this applies to the diabetic kidney or not.

It is showed that in the proximal tubule, GLUT2 translocation to the luminal brush border membrane is related to a rise in facilitative glucose reabsorption (22). The in-vitro research by Ghezzi and Wright, indicated that human SGLT2 was sensitive to insulin and its activity increased significantly by insulin, but human SGLT1 was moderately insensitive to insulin (23). In addition, the in-vitro experiments in proximal tubular cells of rabbits kidney showed that oxidative stress caused by high glucose could reduce the expression and activity of SGLT2 and SGLT1 (24). The inhibition of SGLT2 improves the control of blood glucose in diabetes while the glucose transport increases by SGLT1. However, the quantitative role of kidney SGLT1 in diabetes and kidney damage has not yet been exactly understood, and more studies are needed. Finally the role of sex on benefits of SGLT2 inhibitors in glucose controlling in diabetes is not clear yet, and needs to be specified.

Conclusion

Most findings have revealed a greater proportion of SGLT1 and SGLT2 in female rats compared with males. However in male mice greater SGLT2 existed compared with female mice. However, additional investigations have reported that in humans, these carriers are not affected by gender. In diabetes, SGLT1 expression is reduced in female and male mice and rats. SGLT1 expression in mice and rats reduces, which is to independent gender. To use SGLT inhibitors as therapeutic agents in diabetic condition, the role of sex needs to be more specified.
Authors’ contribution
FK, HHD, SS, MM, NS and MN were involved in preparing the literature review and preparing the draft. The final draft of the article was edited by MN and it was read by all authors. All authors signed the final manuscript.

Conflicts of interest
No conflict of interest was declared by the authors.

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